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Remarks:

This application was filed on 03 - 05 - 2002 as a divisional application to the application mentioned under INID code 62.

(54) Angiostatic steroids

(57) Angiostatic steroids for use in controlling neovascularization and ocular hypertension are disclosed. Pharmaceutical compositions of the angiostatic steroids and mathods for their use in treating neovascularization and coular hypertension, including controlling the ocular hypertension associated with orthany open ande claucoma, are disclosed. In addition, the combination of the compounds with glucocorticoids for the prevention of elevated intraccular pressure during the freatment of inflammation is disclosed.

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#### Description

# Background of the Invention

#### 5 Field of the invention

[0001] This invention relates to angiostatic steroids for controlling coular hypertension. The compounds are also useful in preventing and treating neovascularization. Specifically, the invention is discreted to new angiostatic strates, pharmaceutical compositions comprising the angiostatic steroids, and methods of treatment which comprise administering these compositions to treat coular hypertension, including controlling coular hypertension associated with primary open angle glaucoma, and to freat neovascularization, in addition, the compounds can be used in combination with glucocorticoids to treat coular inflammation without the significant intraocular pressure rise commonly associated with the use of quecoorticoids.

# 5 Description of Related Art

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[D002] Steroids functioning to inhibit angiogenesis in the presence of hepartin or specific heparin fragments are disclosed in Crum, et al., A New Class of Starciels habitate Angiogenesis in the Pressuance of Heparin or a Heparin Fragment, Science, Vol.230, pp.1376-1378 (December 20, 1985). The authors refer to such steroids as "angiostatic" steroids. Included within the new class of steroids found to be angiostatic are the dihydro and tetrahydro metabolities of cortisols and cortexione. In a follow-up study directed to testing a hypothesis as to the mechanism by which the steroids initial angiogenesis, it was shown that heparin/angiostatic steroid compositions cause dissolution of the basement membrane scaffolding to which enchorage dependent endohelik are attached resulting in capillary involution; see, ingior, et al., A Possible Mechanism for Inhibition of Angiogenesis by Angiostatic Steroids: Induction of Capillary Basement Membrane Dissolution, Endocrinology Vol. 119, pp. 1768-1775 19809.

[0003] A group of lartahydro steroids useful in inhibiting angiogenesis is disclosed in international Patent Application No. PCT/US86/02189, Aristoff, et al., (The Ulploin Company). The compounds are disclosed for use in treating head trauma, sepinal trauma, sepine for traumatic shock, stroke and hemorrhage shock, in addition, the patent application discusses the utility of these compounds in embryo implantation and in the treatment of cancer, arthritis and arterioscierosis. Some of the steroids disclosed in Aristoff et al. are disclosed in U.S. Patent No. 4,771,042 in combination with hepsing or a hepsirin fragment for inhibiting angiogenesis in a warm blooded animal.

[0004] Compositions of hydrocortisone, "tetrahydrocortisol-8," and U-72,745G, each in combination with a bata oy-clockwrin, lieve been shown to inhibit comest neovascularization: Li, et al., Angiostatic Steroids Potentistate by Sui-plated Cyclockwrin Inhibit Corneal Neovascularization, Investigative Ophthainology and Visual Science, Vol. 32, No. 11, pp. 2898-2905 (Cotober, 1991). The steroids alone reduce necvescularization somewhat but are not effective alone in effecting redression of neovescularization.

[0005] Tetrahydrocortisol (THF) has been disclosed for its use in lowering the intracoular pressure (10P) of rabbits made hypertensive with dexamethasone/s-beta-dihydrocortisol; see Southren, et al., intracoular Hypotensive Effect of a Topically Applied Cortisol Metabolite: 3-apipla, 5-beta-tetrahydrocortisol, investigate two Ophthalmology and Visual Science, Vol.28 (May, 1987). The authors suggest THF may be useful as an antiglaturoma agent. In U.S. Patern No. 4,863,912, usuand to Southren et al. on September 5,1989, harmaceutical corpositions containing THF and a method for using these compositions to control intracoular pressure are disclosed. THF

has been disclosed as an anglostatic steroid in Folkman, et al., Anglostatic Steroids, Ann. Surg., Vol.206, No.2 (1987) wherein it is suggested anglostatic steroids may have potential use for diseases dominated by shormal necessor-larization, including diabetic ratinopathy, neovascular glaucoma and retrolental fibropiasia. [0008] Many compounds cisselfied as gliucocordicolis, such as discamethasene and prednissionen, are very effective in the treatment of inflammant issues; however, when these compounds are topically applied to the eye to treat coular inflammation, certain patients experience elevated intraocular pressure. Patients who experience these clevations when treated with gliucocordicols are generally interfered to a "steroid' responders." These pressure elevations are of

when tleaned will packcoincoins are guintary interratio to as surrior responders. Treas pressure execution at et or particular concern to pations who already suffer from elevated intracoular pressures, such as galacoma patients. In addition, there is always a nex that the use of glucocorticoids in patients having normal intracoular pressure sew ill cause pressure reser great enough to demage ocular issues. Since glucocorticoid therapy is frequently florg form (le, several days or more), there is potential for eignificant damage to ocular tissue as a result of prolonged elevations in intracoular pressure attributable to that thereav.

[0007] The following articles may be referenced for further background information concerning the well-recognized association between ophthelmic objectorized therapy and elevations in intraccular pressure:

Kitazawa, Increased Intraocular Pressure Induced by Corticosterpids, Am. J. Ophthai., Vol.82 pp.492-493 (1976):

Centrill, et al., Comparison of In Vitro Potency of Corticosteroids with Ability to Raise Intraocular Pressure, Am. J. Ophthal., Vol.79 pp.1012-1016 (1975); and

Mindel, et al., Comparative Ocular Pressure Elevation by Medrysone, Fluorometholone, and Dexamethasone Phosphate, Arch. Ophthal., Vol.98 pp.1577-1578 (1980).

[008] Commonly assigned U.S. Application Sorial No. 07289,351 discloses the use of the angiostatic ateriorit terrahydrocortexolone in combination with a glucocorticold to treat ocular inflammation without the intraooutar pressure elevating effect commonly associated with topical administration of glucocorticolds. In addition, commonly assigned international Application No. PCTI/US90/04071 discloses the angiostatic steroids of Aristoff, et al. In combination with olucocraticolds to veta ocular inflammation without isnificant increase in intraocular pressure.

# Summary of the invention

[0099] This invention is directed to angiostalize steroids and methods of using compositions of these steroids in inhibiting neovascularization. The compositions containing the steroids can be used for treatment of angiogenesis dependent diseases, for example: head fraums, spinal fraums, applie or treumstic shock, stroke, hemorrhagic shock,
cancer, arthritis, arteriosclerosis, angiofibroma, arteriovenous malformations, corneal graft neovascularization, dieseval
wound healing, diebetic retirippethy, granulations, burns, hemangioma, hemophilic joints, hypertrophic scera, necescular glaucoma, nonunion fractures. Osler-Weber Syndrome, psodasis, pyogenic granuloma, retrolental fibropiasia,
poterigium, scleroderma, trachoma, vascular adhesions, and solid tumor growth. In particular, the angiostatic steroids
and compositions hereof are useful for controlling ocular provessoularization.

[0010] The Invention also encompasses methods for controlling ocular hypertension and glaucoma through the systemic or local administration of the compositions disclosed herein.

[0011] The present Invention also includes the use of the angiostatic steroids in combination with glucocorticoids for 5 the treatment of ocular inflammation. The addition of at least one angiostatic steroid makes it possible to employ the potent artificifiammatory glucocorticoids without producing significant delevations in futraccular pressure.

#### Brief Description of the Drawing

[0012] Figure 1 compares the ability of angiostatic steriods to inhibit neovascularization in the rabbit cornea.

#### Detailed Description of Preferred Embodiments

[0013] The development of blood vessels for the purpose of sustaining viable tissue is known as angiogenesis or neovascularization. Agents which inhibit neovascularization are known by a variety of terms such as angiostatic, angiolytic or angiotropic agents. For purposes of this specification, the term "angiostatic agent" means compounds which can be used to control, prevent, or inhibit angiogenesis.

[0014] The angiostatic agents of the present invention are steroids or steroid metabolites. For purposes herein, the term "angiostatic steroids" means steroids and steroid metabolites which inhibit angiogenesis.

[0015] There is currently no effective method for controlling the neovascularization in anglogenesis-dependent diseases. In particular, coular neovascularization has not been successfully treated in the past. Neovascularization of tissues in the front of the eye (i.e. the comes, iris, and the trabecular meshwork) and other conditions, including conditions in the back of the eye, for exemple, retinal, subretinal, macular, and optical nerve head neovascularization, and per prevented and treated by administration of the steroids of the privation are useful in preventing and treating neovascularization, including providing for the regression of neovascularization.

[0018] The anglosiatic steroids can also be used for the control of ocular hypertension, in particular, the agents can be used for the treatment of primary open angle glaucoma.

[0017] The angiostatic steroids of the present invention have the following formula:

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# Structure [A]

# Structure[8]

wherein R<sub>1</sub> is H, β-CH<sub>3</sub> or β-C<sub>2</sub>H<sub>5</sub>;

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Ro is F. Co-Cas double bond, Co-Cas epoxy, H or Cl;

- F<sub>4</sub> is H, CH<sub>3</sub>, Cl or F;
  F<sub>5</sub> is H, OH, F, Cl, Br, CH<sub>3</sub>, phenyl, vinyl or sliyl;

R<sub>R</sub> is H or CH<sub>3</sub>;

 $\mathsf{R}_{3}^{\vee} \ \mathsf{Is} \ \mathsf{CH}_{2} \mathsf{CH}_{2}^{\vee} \mathsf{OR}_{28}, \ \mathsf{CH}_{2} \mathsf{CH}_{2} \mathsf{OC}(=0) \mathsf{R}_{27}, \ \mathsf{H}, \ \mathsf{OH}, \ \mathsf{CH}_{3}, \ \mathsf{F}, = \mathsf{CH}_{2}, \ \mathsf{CH}_{2} \mathsf{C}(=0) \mathsf{OR}_{28}, \ \mathsf{OR}_{28}, \ \mathsf{O}(=0) \mathsf{R}_{27} \ \mathsf{or} \ \mathsf{O}(=0) \mathsf{CH}_{2} \mathsf{C}(=0) \mathsf$ 

35 R<sub>16</sub> is -C=CH, -CH=CH<sub>2</sub>, halogen, CN, N<sub>3</sub>, OR<sub>28</sub>, OC(=O)R<sub>27</sub>, H, OH CH<sub>3</sub> or R<sub>10</sub> forms a second bond between positions C-16 and C-17;

 $R_{12}$  is H or forms a double bond with  $H_1$  or  $R_{14}$ :  $R_{14}$  is Halogen,  $OR_{26}$ ,  $OC(=O)R_{27}$ ,  $NH_2$ ,  $NHR_{26}$ ,  $NHC(=O)R_{27}$ ,  $N(R_{26})_2$ ,  $NC(=O)R_{27}$ ,  $N_3$ , H, -OH, =O, -O-P(=O)(OH)<sub>2</sub>, or -O-C(=O)-(CH)-(CH)-(COOH) where it is an integer from 2 to 6:

40 R<sub>14</sub> is H or forms a double bond with R<sub>12</sub>;

R<sub>15</sub> is H, =O or -OH;

and R<sub>23</sub> with R<sub>10</sub> forms a cyclic phosphate;

wherein Re and Res have the meaning defined above;

or wherein  $R_{23}$  is -OH, O-C(=0)- $R_{11}$ , -OP(0)-(OH)<sub>2</sub>, or -O-C(=0)-(CH<sub>2</sub>)<sub>1</sub>COOH wherein t is an integer from 2 to 6; and  $R_{11}$  is -Y-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-SO<sub>3</sub>H, -Y-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>

wherein Y is a bond or  $\overline{O}$ ; Y' is a bond,  $\overline{O}$ , or  $\overline{O}$ ; each of X and X' is a bond,  $-CON(R_{1g})$ ,  $-N(R_{1g})CO$ , -O, -S, -S,

Z is a bond or -O-; r is an integer of from 2 to 9; and Q is one of the following:

(1) -R<sub>19</sub>: CH<sub>2</sub>COH wherein R<sub>19</sub> is -S<sub>1</sub> -S(O)<sub>1</sub> -S(O)<sub>2</sub> -, SO<sub>2</sub>M(R<sub>20</sub>)-, or N(R<sub>20</sub>)SO<sub>2</sub> ·; and R<sub>20</sub> is hydrogen or lower ally-(C<sub>1</sub>-C<sub>2</sub>); with the proviso that the total number of carbon atoms in R<sub>20</sub> and (CH<sub>2</sub>), is not greater than 10; or (2) -CO-COOH: or

(3) CON(R<sub>21</sub>)CH<sub>2</sub>(R<sub>22</sub>)COOH wherein R<sub>21</sub> is H and R<sub>22</sub> is H, CH<sub>3</sub>, -CH<sub>2</sub>COOH, -CH<sub>2</sub>CH<sub>2</sub>COOH, -CH<sub>2</sub>CH, -CH<sub>2</sub>SH, -CH<sub>2</sub>CH<sub>3</sub>CCH<sub>3</sub>, or -CH<sub>3</sub>Ph-OH wherein Ph-OH is p-hydroxyphenyl;

or R21 is CH3 and R22 is H;

or R2t and R22 taken together are -CH2CH2CH2-;

or -N(R<sub>21</sub>)CH(R<sub>22</sub>)COH taken together is -NHCH<sub>2</sub>COOH; and pharmacoutically acceptable salts thereof; with the proviso that except for the compound wherein R<sub>1</sub> is β-CH<sub>3</sub>, R<sub>2</sub> and R<sub>3</sub> taken together form a double bond between positions 9 and 11, R<sub>4</sub> and R<sub>3</sub> are hydrogen, R<sub>12</sub> and R<sub>3</sub> taken together form a double bond between positions 4 and 5, R<sub>3</sub> is cr.F; R<sub>3</sub> is β-CH<sub>3</sub>, R<sub>10</sub> is cr.OH, R<sub>13</sub> and R<sub>14</sub> taken together form a double bond between positions 4 and 5, R<sub>3</sub> is cr.F; R<sub>3</sub> is β-CH<sub>3</sub>, R<sub>10</sub> is cr.OH, R<sub>13</sub> and R<sub>14</sub> are =0 and R<sub>23</sub> is -OP(O)-(OH)<sub>2</sub>, R<sub>13</sub> is =O only when R<sub>23</sub> with R<sub>3</sub>. Generally the composition of the compo

R<sub>24</sub> = C, C<sub>1</sub>-C<sub>2</sub> double bond, O:

wherein  $R_{26} = C_1 - C_8$  (alkyl, branched alkyl, cycloalkyl, haloalkyl, aralkyl, aryl);  $R_{27} = R_{28} + OR_{28}$ ;  $R_{28} = H$ , C1-C6 (alkyl, branched alkyl, cycloalkyl).

[9018] Excepted from the compounds of Structure [A] are the compounds wherein B<sub>1</sub> is β-CH<sub>3</sub> or β-C<sub>2</sub>H<sub>5</sub>;

R<sub>2</sub> is H or Cl;

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 $R_3$  is  $H_1$ , =0, -OH, O-ality/i(C<sub>1</sub>-C<sub>12</sub>), -OC-i-Olatly/i(C<sub>1</sub>-C<sub>12</sub>), -OC-i-OlAFYL, -OC-OlYi(R), or -OC-OlOF), wherein ARYL is turyl, hieray protyl, or pyridyl and each of said moleties is optionally substituted with one or two (C<sub>1</sub>-C<sub>2</sub>) ality/groups, or ARYL is  $-(CH_3)$ -phenyl wherein if is 0 to 2 and the phenyl ring is optionally substituted with 1 to 3 groups eslected from chlorine, fluorine, bromine, sity/i(C<sub>1</sub>-C<sub>2</sub>), alkoxyi(C<sub>1</sub>-C<sub>3</sub>), thicelikoxy-(C<sub>1</sub>-C<sub>3</sub>), Cig.C<sub>1</sub>, F.5.-, -NHy, and -NHCOCH, and R is hydrocen, that if (C<sub>1</sub>-C<sub>2</sub>), or henvil and each R one the same or different, and - is ARYL as

herein defined, or alkyl( $C_1$ - $C_{12}$ ); or

wherein  $\rm R_2$  and  $\rm R_3$  taken together are oxygen (-0-) bridging positions C-9 and C-11; or wherein  $\rm R_2$  and  $\rm R_3$  taken together form a double bond between positions C-9 and C-11:

or R<sub>o</sub> is α-F and R<sub>o</sub> is β-OH;

or R<sub>2</sub> is α-Cl and R<sub>3</sub> is β-Cl;

and R<sub>4</sub> is H, CH<sub>5</sub>, Cl or F;

Rs is H, OH, F, Cl, Br, CH3, phenyl, vinyl or allyl;

Re is H or CH3;

Fig. is H, OH, CH<sub>3</sub>, F or =CH<sub>2</sub>;

R<sub>10</sub> is H, OH, CH<sub>3</sub> or R<sub>10</sub> forms a second band between positions C-16 and C-17;

R<sub>12</sub> is -H or forms a double bond with R<sub>14</sub>;

R<sub>13</sub> is H<sub>1</sub>-OH, =O, -O-P(O)(OH)<sub>2</sub>, or -O-C(=O)-(CH<sub>2</sub>),COOH where t is an integer from 2 to 6;

R14 is H or forms a double bond with R10:

40 R<sub>15</sub> is =0 or -0H;

and Roa with Ron forms a cyclic phosphate;

wherein Ro and Ros have the meaning defined above:

or wherein R<sub>gs</sub> is -OH, O-C(=0)-R<sub>H</sub>, ... OP(0)-(OH<sub>g</sub>, or -O-C(=0)-(CH<sub>g</sub>)COOH wherein it is an integer from 2 to 8; and R<sub>1</sub> is -Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub>g</sub>)-X,Y-(CH<sub></sub>

Z is a bond or -O-; r is an integer of from 2 to 9; and Q is one of the following:

(1) -R<sub>19</sub>-CH<sub>2</sub>COOH wherein R<sub>19</sub> is -S-, -S(0)<sub>2</sub>-, -S(0)<sub>2</sub>-, -S0<sub>2</sub>N(R<sub>20</sub>)-, or N(R<sub>20</sub>)SO<sub>2</sub>-; and R<sub>20</sub> is hydrogen or lower alkyl-(C<sub>1</sub>-, C<sub>4</sub>); with the proviso that the total number of carbon atoms in R<sub>20</sub> and (CH<sub>2</sub>), is not greater than 10; or minimum and the contraction of the contr

(2) -CO-COOH; or

(a) CON(F<sub>24</sub>)CH(R<sub>22</sub>)COOH wherein R<sub>21</sub> is H and R<sub>22</sub> is H, CH<sub>3</sub>, -CH<sub>2</sub>COOH, -CH<sub>2</sub>CH<sub>2</sub>COOH, -CH<sub>2</sub>OH, -CH<sub>2</sub>OH, -CH<sub>2</sub>CH<sub>3</sub>COOH, -CH<sub>2</sub>OH, -CH<sub>2</sub>OH,

KK

or R<sub>21</sub> is CH<sub>3</sub> and R<sub>22</sub> is H;

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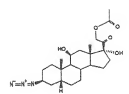
or R21 and R22 taken together are -CH2CH2CH2-

or  $\tilde{\lambda}(R_{p_2})CH(R_{p_2})COOH$  taken togather is  $\tilde{\lambda}$ HiCH\_pCOOH4CH\_pCOOH, and pharmaceutically acceptable salts thereof; with the provisor that except for the compound wherein  $H_1$  is  $\beta$ -CH<sub>3</sub>,  $H_2$  and  $H_3$  taken together form a double bond between positions  $\tilde{s}$  and  $\tilde{H}_1$ .  $H_2$  and  $H_3$  taken together form a double bond between positions  $\tilde{s}$  and  $\tilde{H}_1$ .  $H_3$  is  $H_4$  and  $H_3$  are  $H_4$  and  $H_4$  are  $H_4$  are  $H_4$  are  $H_4$  and  $H_4$  are  $H_4$  are  $H_4$  are  $H_4$  and  $H_4$  are  $H_4$  are  $H_4$  and  $H_4$  are  $H_4$  are  $H_4$  are  $H_4$  and  $H_4$  are  $H_4$  are  $H_4$  and  $H_4$  are  $H_4$  are  $H_4$  and  $H_4$  are  $H_4$  are  $H_4$  are  $H_4$  and  $H_4$  are  $H_4$  are  $H_4$  are  $H_4$  and  $H_4$  are  $H_4$  are  $H_4$  and  $H_4$  are  $H_4$  a

[0019] Unless specified otherwise, all substituent groups attached to the cyclopentanephenanthrene moiety of Structures (A) and (B) may be in either the alpha or bete position. Additionally, the above structures include all pharmaceutically accentible satis of the angiostatic steroids.

[0020] Preferred anglostatic steroids for the treatment of ocular hypertension, neovascular diseases and ocular inllammation are:

38-ACETAMIDO-58-PREGNAN-118,17⊄, 21-TRIOL-20-ONE-21-ACETATE



38-AZIDO-58-PREGNAN-118,17a,21-TRIOL-20-ONE-21-ACETATE

58-PREGNAN-118.17 .. 21-TRIOL-20-ONE

to.

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20-(4-FLUOROPHENYL)THIO-ZI-NOR-5,8-PREGNAN-3~,17~-DIOL

30 20-AZIDO-21-NOR-5\$-PREGNAN-3≪, 17∝-DIOL

20-ACETAMIDO-21-NOR-58-PREGNAN-3«, 17«-DIOL-3-ACETATE

20-(CARBETHOXYMETHYL)THIO-21-NOR-58-PREGNAN-3-,17-DIOL

16 -(2-HYDROXYETHYL)-178-METHYL-58-ANDROSTAM-3=,17=-DIOL

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20-CYANO-21-NOR-5#-PREGNAN-3~,17~-DIOL

17c-METHYL-5β-ANDROSTAN-3c,17β-DIOL

21-NOR-58-PREGN-17(20)-EN-3-CL-3-ACETATE

21-NOR-5#-PREGN-17(20)-EN-3@-OL

21-NOR-5g-PREGN-17(20)-EN-3x-OL-16-ACETIC ACID-3-ACETATE

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21-NOR-5#-PREGNAN-3@,17@,20-TRIOL

21-NOR-58-PREGNAN-17=,20-DIOL-3-ACETATE

4,9(11)-PREGNADIEN-17×,21-DIOL-3,20-DIONE-21-ACETATE

4,9(11)-PREGNADIEN-17∝,21-DIOL-3,20-DIONE

11-EPICORTISOL

17 - HYDROXYPROGESTERONE

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TETRAHYDROCORTEXOLONE (THS)

TETRAHYDROCORTISOL (THF)

21-HOR-58-PREGN-17(20)-EN-34, 16-DIOL-3-ACETATE-16-(0-METHYL)MALONATE 21-MOR-5~-PREGNAN-3«,17«,20-TRIOL-3-PHOSPHATE

21-NOR-5#-PREGN-17(20)-EN-3m,16-DIOL

21-NOR-58-PREGNAN-3=,178,20-TRIOL

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21-NOR-5x-PREGNAM-3x,178,20-TRIOL

4-ANDROSTEN-3-DNE-178-CARBOXYLIC ACID

OH -

17∝-ETHYNYL-5(10)-ESTREN-17#-OL-3-ONE

17-ETHYMYL-1,3,5(10)-ESTRATRIEN-3,17#-DIOL

[0021] Most preferred compounds for preventing and treating neovascularization are:

4.9(11)-Pregnadien-17a,21-dioi-3,20-dione-21-acetate

21-Nor-5β-pregn-17(20)-en-3α,16-diol-3-acetate-16-(O-methyl)malonate

4,9(11)-Pregnadien-17a,21-diol-3,20-dlone

21-Nor-5β-pregnan-3α,17α,20-triol-3-acetate

21-Nor-5α-pregnan-3α,17α,20-triol-3-phosphate

[0022] The angiostatic steroids of the present invention are useful in inhibiting neovascularization and can be used in treating the neovascularization associated with: head trauma, spinal trauma, systemic or traumatic shock, stroke, hemorrhagic shock, cancer, artifitis, arterioscenous, angiofitroma, arteriovenous malformations, comeal graft neovascularization, delayed wound healing, diabetic retinopathy, granulations, burns, hemangioma, hemophilic joints, hypertrophic scars, neovascular jaucorins, nonunion fractures, Osler-Weber Syntrome, psorlasis, properlic granuloma, retroehant libropalish, betrafficium, scleroderma; trachoma, vascular adhesions, and solid tumor growth,

[0023] In particular, the angiostatic steroids are useful in preventing and freeling any ocular necvascularization, including, but not limited to: retinal diseases (diabetic retinopathy, chronic glaucoma, rathal detachment, sickle cell retinopathy, serille macular degeneration due to subretinal neovascularization); rubecels iritis; inflammatory diseases; chronic uveitis; neoplasms (retinoblastoma, pseudoglioma); Fuchs' heterochromic iridocyclitis; neovascularization (inflammatory, transplantation, developmental hypoplasia of the Irity); neovascularization fullmamatory, transplantation, developmental hypoplasia of the Irity; neovascularization fullmamatory, transplantation, developmental hypoplasia of the Irity; neovascularization resulting following a combined vitrectomy and lensectomy; vascular diseases (retinal ischemia, choroidal vascular insufficiency, choroidal thrombosis, carotid artery ischemia); plerigium; neovascularization of the eye or confusive coular injury.

[00:24] The initiation of new blood vessel formation may arise quite differently in various tissues or as a result of different diseases. Many substances have been found to induce necessualization, see, Folkman, et al., Anglogenic Factors, Science, Volume 235, pp. 442-447 (1987), However, It is believed, that once initiated, the process of neovascularization is similar in all tissues regardless of the associated disease, Furth, Critical Factors Controlling Analogen-

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esis. Cell Products, Cell Metrix, and Growth Factors, Laboratory Investigation, Volume 55, No. 5, pp. 505-609 (1986). [0025] There are a variety of theories regarding the mechanism of section of angiostatic steroids. For example, angiostatic steroid induced inhibition of neovascularization may occur due to, dissolution of the expiliary basement mambrane, Inglane, et al., Supre; inhibition of viaurate rendtheisal cell profileration. Cartiou, et al., Inhibition of viaurate rendtheisal cell profileration. Cartiou, et al., Inhibition of viaurate rendtheisal cell profileration. Cartiou, et al., Inhibition of viaurate rendtheisal cell profileration. The profileration by Heparin and Steroids, Cell Biology International Reports, Vol. 12, No. 12, pp. 1037-1047 (December, 1988); effect on vascular endotheisal cell laminini expression, Tokida, et al., Production of Tieu Viainal Laminini Forms by Endotheisal Cells and Still of Their Relative Levels by Angiostatic Staroids, The Journal of Tieu Viainal Laminini Forms by Endotheisal Cells and Still of Their Relative Levels by Angiostatic Staroids, The Journal of Tieu Viainal Laminini Forms et al., Antiangiopenic Action of Heparin Plus Corticone is Associated with Decreased Collagenous Synthesis, Mariana and Cartious Collagenous Staroids in the Chick Chortosillantoic Membrane System, The Journal of Phermacology and Experimental Therapoutics, Vol. 251, No. 2, pp. 519-828 (1989); and inhibition of vascular endotheisal cell plasminopen activator extity, Ashitor-Fuse, et al., Medroxypropesterone Acetate, An Anti-Cancer and Anti-Angiogenic Steroid, finibits the Plasminopen Activator in Bowine Endotheisal Colls, Int. J. Cancer, 44, pp. 859-884 (1989).

[0026] There are many theories associated with the cause of neovascularization, and there may be different inducers depending on the disease or surgery involved, BenEzra, Neovasculogenic Ability of Prostaglandins, Growth Factors, and Synthetic Chemoattractants, American Journal of Ophthalmology, Volume 86, No. 4, pp. 455-461, (October, 1978). Regardless of the cause or the associated disease or surgery, it is believed that angiostatic agents work by inhibiting one or more steps in the process of neovascularization. Therefore, the angiostatic steroids of this invention are useful in the treatment and prevention of neovascularization associated with a variety of diseases and surgical complications. [0027] The anglostatic steroids of the present invention may be incorporated in various formulations for delivery. The type of formulation (topical or systemic) will depend on the site of disease and its severity. For administration to the eve, topical formulations can be used and can include ophthalmologically acceptable preservatives, surfactants, viscosity enhancers, buffers, sodium chioride, and water to form aqueous sterile ophthalmic solutions and suspensions. in order to prepare sterile orbithalmic ointment formulations, an angiostatic steroid is combined with a preservative in an appropriate vehicle, such as mineral off, liquid lanolin, or white petrolatum. Sterile ophthalmic gel formulations comprising the anglostatic steroids of the present invention can be prepared by suspending an anglostatic steroid in a hydrophilic base prepared from a combination of, for example, Carbopol® (a carboxy viny) polymer available from the BF Goodrich Company) according to published formulations for analogous ophthalmic preparations. Preservatives and antimicrobial agents may also be incorporated in such gel formulations. Systemic formulations for treating ocular ne-

injection.

[0028] The specific type of formulation selected will depend on various factors, such as the anglostatic steroid or its salt being used, the dosage frequency, and the location of the necvascularization being treated. Topical ophthalmic accuracy and continued to the continued of the second of the back of the sey if the angiostatic agent can be formulated such that it can be delivered topically and the agent is able to penatrate the issues in the front of the eye. The angiostatic steroid will normally be contained in these formulations in an amount from about 0.01 to about 1.0 a weight/percent. Preferable concentrations range from about 0.1 to about 5.0 weight/percent. Thus, for topical administration, these formulations are delivered to the surface of the eye one to six times a day, depending on the routine discretion of the skilled clinician Systemic administration, for example, in the form of tablets is useful for the treatment of neovascularization particularly of the back of the eye, for example, the rental. Tablets containing 10-100 mg of anglostatic agent can be taken 2-3 times per day depending on the form of the discretion of the skilled clinician services and the surface of the discretion of the skilled clinician services and maintains of the services of the servi

ovascularization can also be used, for example, orally ingested tablets and formulations for intraocular and periocular

above act to centrol intracoular pressure by inhibiting the accumulation or stimulating the dissolution of smorphous stracellular material in the Itabecular meshwork of the eye. The presence of this amorphous extracellular material atters the integrity of the healthy trabocular meshwork and is a symptom associated with primary open angle glaucoma (PCAG), it is not well understood why this amorphous extracellular material builds up in the trabocular meshwork of persons suffering from PCAG. However, it has been found that the amorphous extracellular material is generally composed of glycosaminophycans (GAGS) and basement membrane material; see, Ophthalmology, Vol.30, No.7 (July 1983); Mayor Carlo, Proc. Vol.51, pp.50-97 (Jun. 1985); and Potalic Neurosci. Vol.12, pp.240-251 (1985-38), Winnor these materials build up in the trabocular meshwork, the squeous humor, normally present in the anterior charmor of the eye, cannot leave this chamber through its normal route (the trabocular meshwork) at its normal rate. Therefore, a normal volume of aqueous humor is produced by the ciliary processes of the eye and introduced into the anterior chamber, but its exit through the trabocular meshwork is also normally solven. This results in a buildup of pressure in the shadors.

skilled clinician.

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eye, ocular hypertension, which can translate into pressure on the optic nerve. The ocular hypertension so generated can lead to blindness due to damage to the optic nerve.

[0031] Many methods for treating primary open angle glaucoma and ocular hypertension concentrate on the production of aquious humor by the eye, thowever, aquious humor is the fundamental source of nourishment for the issues of the eye, particularly the cornea and lens which are not sustained by blood supply. Therefore, it is not desirable to deprive these lissues of the necessary ingigation and nutrifion provided by the aquious humor. It is desirable to the tor normal exit of the aquious humor, it is desirable to the tor normal exit of the aquious humor, it is desirable to the tor normal exit of the aquious humor, the substantial interest in the normal integrity of the trebecular meshwork. This is accomplished according to the present invention by the administration of anniciating is steriols.

[0032] It is believed that the angiostatic steroids disclosed berein function in the trabecular meshwork in a similar manner as shown by righer, et al., wherein it was shown that angiostatic steroids caused disclosulino of the basement membrane scaffolding using a chick embryo neovascularization model; Endocrinology, 119, pp.1768-1775 (1988), it is believed that the angiostatic steroids of the present invention prevent the accumulation, or promote the dissolution of, amorphous extracollular materials in the trabecular membrane with province of the formation of besement membrane materials and glycosarvinoplycans. Thus, by preventing the development of these materials or promoting their dissolution, the normal infective of the trabecular membrane; and and success humor may flow through the trabecular membrane; and and success humor may flow through the trabecular membrane; and and success humor may flow through the trabecular membrane; and and success humor may flow through the trabecular membrane; and and success humor may flow through the trabecular membrane; and and success humor may flow through the trabecular membrane; and and success humor may flow through the trabecular membrane; and and success humor may flow through the trabecular membrane; and and success the success of the state of the success of the state of t

meshwork at normal rates. As a result, the intraocular pressure of the eve is controlled.

10033] The anglostatic steroids of the present invortion may be incorporated in various formulations for delivery to the eye to control coular hypertension. For example, topical formulations can be used and can include ophthalmosphologically acceptable preservatives, surfactants, viscosity enhancers, buffers, sodium chloride and water to form aqueous sterile ophthalmic oliment formulations, an angiostatic steroid is combined with a preservative in en appropriate veribide, such as mineral oil, figuid familion of white periodistrum. Sterile ophthalmic gel formulations compressing the angiostatic steroids of the present invortion can be prepared by suspending an angiostatic steroid in a hydrophilio base prepared form a combination of, for example, Carbopol@404 (a carboxyvinyl polymer evallable from the B.F. Goodrich Company) according to published formylations or analogous ophthalmic preparations. Preservatives and tonicity agents may also be incorporated in such gli formulations. The specific type of formulations selected will depend on various factors, such as the angiostatic steroid or its satishing used, and the dosage frequency. Topical ophthalmic apressed frequency. Topical ophthalmic apressed frequency inspiral ophthalmics appeared in such gale are the preferred dosage forms. The angiostatic steroid will normally be contained in these formulations in an amount of from about 0.005 to about 5.0 weight precent (wf.5). Preferebel concentrations range from sebout 0.05 to about 5.0 weight precent (wf.5). Preferebel concentrations range from sebout 0.00 to be bout 5.0 weight precent the skilled clinicions are delivered to the surface of the eye one to four times per daw depending upon the proferred obsequence of the skilled clinicions are delivered to the surface of the eye one to four times per daw depending upon the proferred obsequence of the skilled clinicions are delivered to the surface of the eye one to four times per daw depending upon the proferred to the surface of the eye one to four times per daw depen

[0034] In addition, artilinflammatory compositions of glucocorticoids can contain one or more anglostatic steroids of the present invention, preferably tetrahydrocortisol. These compositions will contain one or more anglostatic steroids of the present invention in an artilinflammatory effective amount and will contain one or more anglostatic steroids of the present invention in an amount effective to inhibit the IOP elevating effect of the glucocorticoids. The amount of each component will depend on various factors, such as the reliably endemoty of certain glucocorticoids to ususe IOP elevations, the servity and type of ocular inflammation being treated, the estimated duration of the treatment, and so on. In general, the ratio of the amount of glucocorticoid to the amount of anglostatic steroid on a weight to weight basis will be in the range of 10: 1 to 1:20. The concentration of the glucocorticoid component will typically be in the range of about 0.01% to about 5.0% by weight. The concentration of the angiostatic steroid component will typically be in the range of about 0.05% to about 5.0% by weight.

[0035] The above-described active ingredients may be incorporated into various types of systemic and ophthalmic inemutations. For example, for topical coular administration, in the active ingredients may be combined with ophthalmologically acceptable preservatives, surfactants, viscosity enhancers, buffers, toxicity agents and water to form an aquestic state of the properties of the properties are combined with a preservative in an appropriate vehicle, such as mineral oil, liquid lanolin, or white petrolatum. Sterile ophthalmic ophthalmic pole formulations may be prepared by suspending the active ingredient in a hydrophilic base prepared from the combination of Carbopol \$40 (a carboxy vinly polymer available from the E.F. Goodrich Company) according to published formulations for analogous ophthalmic preparations; preservatives and tonicity agents can also be incorporated. The specific type of formulation solected will depend on various factors, such as the severity and type of ophthalmic inflammation being treated, and desage frequency. Ophthalmic solutions, suspensions, oitments and gelies are the preferred dosage forms, and topical spoilcation to the inflammaduling the specific type of formulation seleptication to the inflammaduling the specific type of proprietd on the general dosage forms, and topical spoilcation to the inflammaduling the specific type of proprietd of the general dosage forms, and topical spoilcation to the inflammaduling the propriet of the specific type of proprietd of the general dosage forms, and topical spoilcation to the inflammaduling the specific type of the preferred dosage forms, and topical spoilcation to the inflammaduling the propriet of the propri

[0036] The following examples illustrate formulations and synthesis of compounds of the present invention, but are in no way limiting.

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# Example 1

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[0037] The topical compositions are useful for controlling ocular hypertension or controlling ocular neovasculariza-

Component	wt.%
Angiostatic Steroid	0.005-5.0
Tyloxapol	0.01-0.05
НРМС	0.5
Benzalkonium Chloride	0.01
Sodium Chloride	0.8
Edetate Disodium	0.01
NaOH/HCI	q.s. pH 7.4
Purified Water	q.s. 100 mL

# Example 2

[9038] The composition is useful for controlling ocular hypertension.

Component	wt.%
21-Nor-5β-pregnan-3α,17α,20-tricl	1.0
Tylexapol	0.01-0.05
HPMC	0.5
Benzalkonium Chloride	0.01
Sodium Chloride	0.8
Edetate Disodium	0.01
NaOH/HCI	q.s. pH 7.4
Purified Water	q.s. 100 mL

- [0039] The above formulation is prepared by first placing a portion of the purified water into a beaker and heating to 90°C. The hydroxypropylmethylocilulose (HPMC) is then added to the heated water and mixed by means of vilgorous vortex stirring until all of the HPMC is dispersed. The resulting mixture is then allowed to pool while undergoing mixing in order to hydrate the HPMC. The resulting solution is then sterilized by means of autoclaving in a vessel having a liquid inlet and a hydroxhobic, sterile air vent filter.
- [0040] The sodium chloride and the edetate disodium are then added to a second portion of the purified water and dissolved. The benzalkonium chloride is then added to the solution, and the pH of the solution is adjusted to 7.4 with 0.1 M NaCH/HCI. The solution is then sterilized by means of filtration.
- [0041] 21-Nor-5β-pregnan-3cu,17c,20-triol is sterifized by either dry heat or ethylene oxide. If ethylene oxide sterifization is selected, aeration for at least 72 hours at 50°C, is necessary. The sterifized steroid is weighed aseptically and placed into a pressurized ballmill container. The tytoxapol, in sterifized aqueous solution form, is then added to the ballmill container. Sterifized glass balls are then added to the container and the contents of the container are milled assptically at 225 pm for 16 hours, or until all particles are in the range of approximately 6 microns.
- [0042] Under asspite conditions, the micronized drug suspension formed by means of the preceding step is then poured into the HPMC solution with mixing. The ballmill container and balls contained therein are then rinsed with a portion of the solution containing the sodium chloride, the detatar discribum and benzalkonium chloride. The rinse is then added aseptically to the HPMC solution. The final volume of the solution is then adjusted with purified water and, if necessars, the BH of the solution is adjusted to the TA with NoOHHPIC.

# Example 3

19943] The following formulation is representative of the antiinflammatory compositions of the present invention.

Component	wt.%
4.9(11)Pregnadien-17α,21-diol-3,20-dione-21-acetate	1.0
Dexamethasone	0.1
Tyloxepol	0.01 to 0.05
HPMC	0.5
Benzalkonium Chloride	0.01
Sodium Chloride	0.8
Edetate Disodium	0.01
NaOH/HC!	q.s. pH 7.4
Purified Water	q.s. 100 mL

[9044] The above formulation is prepared in the same manner set forth in Example 2, sterilizing and adding the dexamethasone to the steroid before placing both into a pressurized ballmill container.

# Example 4

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[0045] The following formulation is another example of the antiinflammatory compositions of the present invention

	wl.%
Tetrahydrocortisol	1.0
Prednisolone Acetate	1.0
Tyloxapol	0.01 to 0.05
HPMC	0.5
Benzaikonium Chloride	0.01
Sodium Chloride	0.8
Edetate Disodium	0.01
NaOH/HCI	q.s. pH 7.4
Purified Water	q.s. 100 mis

<sup>36</sup> [0046] The above formulation is prepared in the same manner set forth in Example 2, sterilizing and adding the pradmiscione acetate to the sterild before placing both into a pressurized belimili container. [0047]. The following formulations are representative of compositions used for the treatment of angiogenesis dependent diseases.

# 40 Example 5

FORMULATION FOR ORAL ADMINISTRATION

# [0048]

Tablet

10-1000 mg of anglostatic steroid with inactive ingredients such as starch, lactose and magnesium stearate can be formulated according to procedures known to those skilled in the art of tablet formulation.

# 50 Example 6

FORMULATION FOR STERILE INTRAOCULAR INJECTION

# [0049]

each mL contains:	
Angiostatic Steroid	10-100 mg
Sodium Chloride	7.14 mg
Potassium Chloride	0.38 mg
Calcium chloride dihydrate	0.154 mg
Magnesium chloride hexahydrate	0.2 mg
Oried sodium phosphate	0.42 mg
Sodium bicarbonate	2.1 mg
Dextrose	0.92 mg
Hydrochloric acid or sodium hydroxide to adjust pH to approximately 7.2 Water for Injection	

6 Example 7

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FORMULATION FOR TOPICAL OCULAR SOLUTION

20 [0050]

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21-Nor-5α-pregnan-3α,17α-20-triol-3-phosphate	1.0%
Benzalkonium chloride	0.01%
HPMC	0.5%
Sodium chloride	0.8%
Sodium phosphate	0.28%
Edetate disodium	0.01%
NaOH/HCI	q.s. pH 7.2
Purified Water	q.s. 100 mL

# Example 8

FORMULATION FOR TOPICAL OCULAR SUSPENSION

# [0051]

Ingredient	Amount (wt.%)
4,9(11)-Pregnadien-17α,21-diol-3,20-dione-21-acetate	1.0
Tyloxapol	0.01 to 0.05
HPMC	0.5
Benzalkonium chloride	0.01
Sodium chloride	0.8
Edetate Disodium	0.01
NaOH/HCI	q.s. pH 7.4
Purified Water	q.s. 100 mL

- [0052] The formulation is prepared by first placing a portion of the purified water into a beaker and heating to 90°C. The hydroxypropyimethy/cellulose (HPMC) is then added to the heated water and mixed by means of vigorous votex stirring until all of the HPMC is dispersed. The resulting mixture is then allowed to cool while undergoing mixing in order to hydrate the HPMC. The resulting solution is then sterifized by means of autoclaving in a vessel having a liquid intel and an hydrophobic, sterified air ventifilter.
- [0053] The socium chloride and the edelate disodium are then added to a second portion of the purified water and dissolved. The benzalkonium chloride is then added to the solution, and the pH of the solution is adjusted to 7.4 with 0.1 th NaO/PHCI. The solution is then stellized by means of filtration.
  - [9954] The 4,9(11)-Pregnadien-17a,21-dioi-3,20-dione-21-acetate is sterilized by either dry heat or ethylene oxide.

If othlyene oxide sterilization is selected, aeration for at least 72 hours at 50°C is necessary. The sterilized 4,9(11) — Pregnandien-17a,21-dipi-3,20-dione-21-acetate is weighed aseptically and placed into a pressurized ballmill container. The tyloxappi, in sterilized aqueous solution form, is then added to the ballmill container. Sterilized gleas balls are then added to the container and the contents of the container are milled asoptically at 225° pm for 16 hours, or until all particles are in the range of approximately 6 microns.

[0055] Under aseptic conditions, the micronized drug suspension formed by means of the preceding step is then poured into the HPMC solution with mixing. The ballmill container and balls contained threein are then rinsed with a portion of the solution containing the sodium chloride, the edetate disodium and benzalkonium chloride. The rinse is then added asspeciately to the HPMC solution. The final volume of the solution is then adjusted with purified water and, if necessary, the pH of the solution is adjusted to pH 7.4 with NaCHPHC. The formulation will be given topically, in a therapeutically effective amount, in this instance, the phrase "therapeutically effective amount" means an amount which is sufficient to substantially prevent or reverse any coular proviscoularization. The dosage regimen used will depend on the nature of the neovescularization, as well as various other factors such as the patient's age, sex, weight, and medical history.

# Example 9

FORMULATION FOR OBAL ADMINISTRATION

#### 20 100561

Tahle

5-100 mg 21-Nor-5β-pregnan-3α-17α-20-triol with inactive ingredients such as starch, factose and magnesium stearate can be formulated according to procedures known to those skilled in the art of tablet formulation.

#### Example 10

Formulation for Sterile Intraocular Injection

#### 30 [0057]

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each mL contains:	
4,9(11)-Pregnadien-17α,21-diol-3,20-dione	10-100 mg
Sodium Chloride	7.14 mg
Potassium Chloride	0.38 mg
Calcium chloride dihydrate	0.154 mg
Magnesium chloride hexahydrate	0.2 mg
Dried sodium phosphate	0.42 mg
Sodium bicarbonate	2.1 mg
Dextrose	0.92 mg
Hydrochloric acid or sodium hydroxide	
to adjust pH to approximately 7.2	
Water for injection	

# Example 11

inhibition of angiogenesis in the rabbit corneal neovascularization model:

[0058] The corneal pocket system of BenEzra (Am. J. Ophthalmol 88.455-461, 1978) was used to induce corneal neavescularization in the rabible. A amell Enwar pellet containing 0.54g of lipophysaccharide (LPS) was lineared into the middle of the corneal stroma and positioned 2.5 mm from the limbus. An additional Elvax pellet with or without 50 gg of angiosiatic storiol was placed next to the LPS implant. The eyes were examined daily and the area of neovascularization calculated. Results after 8 days of LPS implantation are snown in Figure 1. The \*-tetrahydrocorists\* (A = 4.9(11)-Pregnadien-17a;21-dioi-3.20-dione-21-accitate; B = 4.9(11)-pregnadien-17a;21-dioi-3.20-dione-21-accitate;

# Example 12

# Preparation of 5β-Pregnan-11β, 17α, 21-triol-20-one

# Tetrahydrocortisoi-F-21-t-butyldiphenylsilyi ether (PS03842)

[0059] A solution of 4.75 g (17.9 mma) of t-buyldiphenylchionosilane in 5 mL of dry DMF was added dropwase to a strines deution of 5.7 g (15.6 mmo) of tetrahylchocortisol-F (Steratoldis N.P. 9050) and 2.3 g (15 mma) of 4-dimber to a strines obtained (DMAP) in 30 m L of dry DMF, under N<sub>2</sub>, at .25 to .30°C (maintained with CO<sub>2</sub> - MeCN). After a further 20 min at .30°C. the mbuture was allowed to warm to 25°C overnicith.

[9060] The mixture was partitioned between ether and water, and the organic solution was washed with brine, dried

(MgSO<sub>4</sub>), filtered and concentrated to give 10.7 g of a white foam,

100613 This material was purified by flash column chromatography (400 g silice; 82.5 to 79% ether/hoxane). The 3-elioxy isomer eluted first, followed by mixed fractions, followed by the little compound. The concentrated mixed fractions (4.0 g) were chromatographic on the same column with 35% ethyl acettate/hexane. The total yield of the 3-elioxy isomer was 0.42 g (5%), and of the title compound, 5.05 g (53.5%). Continued elution with 25% MeOH/EIOAc allowed recovery of unreacted tetra/hoxocratio-1F.

# PS03842

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[0062] NMR (200 MHz ¹+l) (CDCl<sub>3</sub>): 80.63 (s. 3H, Me-18); 1.11 (s. 9H, I-Bu); 1.12 (s. 3H, Me-19); 2.57 (t, J=13, 1H, I+B); 2.6 (s. 1H, CH-17); 3.63 (sept, J-2.5, 1H, I+3); 4.15 (br.s., 1H, I+11); 4.37 and 4.75 (AB, J=20, 2H, I+21); 7.4 (m. 8H) and 7.7 (m. 4H) (Ph.).

NMR (200 MHz <sup>1</sup>H) (DMSO-3<sub>8</sub>); 80.84 (s, 3H. Me-18); 1.02 (s, 9H, t-Bu); 1.07 (s, 3H, Me-19); 2.50 (t, J=13, 1H, H-8); 3.37 (m, 1H, H-3); 3.94 (d, J=2, 1H, OH-11); 4.00 (br s, 1H, H-11); 4.42 (d, J=5, 1H, OH-3); 4.38 and 4.83 (AB, J=20,

2H, H-21); 5.11 (s, 1H, OH-17); 7.45 (m, 6H) and 7.8 (m, 4H) (Phg).

[0083] NMR (60.3 - MHz <sup>16</sup>0) (CDCl<sub>2</sub>): 17.4 (C-18): 19.3 (C-16); 23.7 (C-15): 25.8 (C-7): 25.8 (C-7): 25.8 (Ms<sub>2</sub>C): 27.2 (C-6): 30.9 (C-2): 31.5 (C-6): 34.1 (Ms<sub>2</sub>C): 34.8 (C-10): 35.2 (C-1): 36.2 (C-4): 39.7 (C-13): 35.5 (C-5): 44.3 (C-6): 37.4 (C-12): 52.1 (C-14): 67.8 (C-11): 68.9 (C-21): 71.7 (C-2): 83.8 (C-14): 127.8 (127

[0064] [R (KBr) 3460, 2930, 2860, 1720, 1428, 1136, 1113, 1070, 1039, 703 cm<sup>-1</sup>.

[0065] This compound did not show a sharp melting point but turned to a foam at 80-100°C. Numerous attempts at recrystallization failed.

#### 58-Pregnan-116, 17a, 21-triol-20-one

[0068] A solution of PS03842 (0.91 g, 1.50 mmol) and thiocarbonyl dilmidazole (1.05 g, 5.9 mmol) in 8 mL of anhydrous dioxane was refluxed under N<sub>2</sub> for 3.5 h. The cooled solution was partitioned between either and water and the organic solution was washed with brine, dried (MgSQ<sub>4</sub>), filtered and concentrated. The residue was chromatographed (120 g SiO<sub>2</sub>, 35% EiOA2/mexane) giving 0.86 g (80%) of the imidazolyl thiosetter.

[0067] A solution of 0.75 g (1.05 mmol) of this compound in 100 mL of anhydrous dioxane was added dropwise over 2.2 h to a rapidly stirred, raffuxing solution of 1.6 mL (5.9 mmol) of BugSnH in 100 mL of anhydrous dioxane unifor N<sub>2</sub>. After a further 1 h at raffux, the solution was cooled, concentrated and the residue chromatographete (200 g STG<sub>2</sub>, 9% EiCAchexane) giving 0.43 g (70%) of the 3-deoxy-21-silyl ether. This material was dissolved in 20 mL of methanol;

ECOAChexane) giving 0.43 g (70%) of the 3-deoxy-21-skilly ethick. This material was dissolved in 2 or 10 of mismanol SigNF-64P, 0.65 g 1.8 mind) was added, and the mixture was heated to reflux under N<sub>2</sub> for 4 h. The cooled solution was diffused with 2 volumes of EtOAc, concentrated to 1/4 volume, partitioned (EtOAc/H<sub>2</sub>O), and the organic solution was washed with brine, dried (MgSO<sub>4</sub>), filtered and concentrated. The residue (0.40 g) was chromatographed (30 g SiO<sub>2</sub>, 40% EtOAchexane) to time 0.25 g (98%) of an oil.

100683 This oil was crystallized (n-BuCl to afford 0.14 g of the title compound as a white solid, m.p. 167-170°C.

100691 IR (KBr): 3413 (br), 2934, 1714, 1455, 1389, 1095, 1035 cm<sup>-1</sup>,

f00701 MS (CI): 351 (M+1).

[0071] NMR (200 MHz <sup>1</sup>H, DMSO-d<sub>e</sub>); 80.89 (s, 3H, Me-18); 1.14 (s, 3H, Me-19); 0.8-2.0 (m); 2.5 (t, J=13, 1H, H-8), 3.96 (d, J=2, 1H, OH-11); 4.1 (br.s, 1H, H-11); 4.1 and 4.5 (AB, Jurther split by 5 Hz, 2H, H-21); 4.6 (t, J=5, 1H, OH-21); 5.14 (s, 1H, OH-17).

[0072] Arial. Calc'd for C<sub>21</sub>H<sub>34</sub>O<sub>4</sub>: C, 71.96; H, 9.78. Found: C, 71.69; H, 9.66.

#### Example 13

# Preparation of 21-Methyl-5β-pregnan-3α, 11β, 17α, 21-tetrol-20-one 21-methyl ether

[9973] Sodium hydride (60% cil dispersion, 9,10 g, 2.5 mmol) was added to a stirred solution of tetrahydrocortisol-F (0.73 g, 2.0 mmol) and CH<sub>3</sub>I (0.60 mL, 9.6 mmol) in 8 mL of anhydrous DMF under No. Hydrogen was evolved, and the temperature rose to 35°C. After 1 h, the mixture was diluted with EtOAc, extracted with water (until neutral) and brine, dried (MgSO<sub>4</sub>), filtered and concentrated. The residue was chromatographed (70 g SiO<sub>5</sub>, 80% EIOAc/hexane) to give 0.17 g of a white solid, MS (CI) = 395 (M +1). This material was recrystallized (EtOAc-n-BuCI) to afford 0.12 g

(16%) of the title compound as a feathery white solid, m.p. 208-213 °C.

[9074] IR (KBr): 3530, 3452, 2939, 2868, 1696 (s. CO), 1456, 1366, 1049 cm<sup>-1</sup>.

[10075] NMR (200 MHz 1 H, DMSO-de); 80.74 (s. 3H, Me-18); 1.09 (s. 3H, Me-19); 1.14 (d. J=6.6, 3H, C-21 Me); 9.8-2.0 (m): 2.47 (t, J=13, 1H, H-8): 3.18 (s, 3H, OMe): 3.35 (m, 1H, H-3): 4.00 (d, J=2, 1H, OH-11): 4.07 (br s, 1H, H-11); 4.37 (g, J=6.8, 1H, H-21); 4.43 (d, J=5, 1H, OH-3); 5.16 (s, 1H, OH-17).

[0076] Anal. Calc'd for CoallingOs: C, 70.01; H, 9.71.

Found: C, 70.06; H, 9.76.

# Example 14

# Preparation of 38-Azido-56-pregnan-118, 17g, 21-triol-20-one-21-acetate

[9077] A solution of triphenylphosphine (2.6 g., 10 mmol) in 10 mL of toluene was carefully added to a stirred solution of PS03842 (see Example 4) (1.75 g, 2.90 mmol), diphenylphosphoryl azide (2.2 mL, 10.2 mmol) and diethyl azodicarboxylate (1.55 mL, 10 mmol) under N<sub>2</sub>, keeping the internal temperature below 35°C (exothermic). The solution was stirred for 1.2 h, then diluted with ether, washed with water and brine, dried (MgSQ<sub>4</sub>), filtered and concentrated and the residue (9.5 g. oil) chromatographed 175 g SiO2, 15% EtOAc/hexane) giving 1.83 g of a viscous oil. [9078] A solution of 1.73 g of this material and 1.75 g (5.5 mmol) of 8u NF-3H<sub>2</sub>O in 20 mL of methanol was refluxed under No for 2.5 h. The crude product (1.94 g) was isolated with ethyl acetate and chromatographed (100 g SiOn, 50% EtOAc/hexane) giving 0.60 g (55%) of a white semisoild. Trituration (4:1 hexane-ether) gave 0.57 g (53%) of a solid. [0079] A stirred solution of 0.40 g of this material in 3 mL of dry pyridine was treated with 0.3 mL of acetic anhydride and stirred overnight at 23°C under No. The mixture was quenched with 1 mL of methanol, stirred for 15 min, diluted with ether, washed with 1 M aqueous HCI, water (until neutral), brine, dried (MqSQ4), filtered and concentrated. The residue (0.41 g, oil) was chrometographed (35 g SiOs, 33% EtOAc/hexane) to afford 0.33 g (76%) of the title compound as a white foam, m.p. 80-90°C (dec).

[0080] IR (KBr): 3505, 2927, 2866, 2103 (vs), 1721 (sh 1730), 1268, 1235 cm<sup>-1</sup>.

[0081] NMR (200 MHz 1H, CDCl<sub>3</sub>): 80.92 (s, 3H, Me-18); 1.21 (s, 3H, Me-19); 1.0-2.1 (m); 2.17 (s, 3H, Ac); 2.26 (s 1H, OH-17); 2.74 (m, 1H, H-8); 3.97 (br s, 1H, H-3); 4.31 (br s, 1H, H-11); 4.94 (AB, J=17, Av=60, 2H, H-21). [0082] Anal. Calc'd for C23H35N3O5: C, 63.72; H, 8.14; N, 9.69.

Found: C, 63.39; H, 8.18; N, 9.45.

# Example 15

# Preparation of 3β-Acetamido-5β-pregnan-11β, 17α-21-triol-20-one-21-acetate

- [0083] A solution of 3β-azido-5β-pregnan-11β,17α,21-triol-20-one-21-acetate (0.15 g, 0.35 mmol) in 8 mL of absolute ethanol containing 0.03 g of 10% Pd on C was stirred under Ho (1 atm) at 23°C for 2 h, The mixture was filtered and concentrated, the residue dissolved in EtOAc, the basic meterial extracted into 1 M aqueous HCI, liberated (Na-CO<sub>2</sub>), extracted (EtOAc) and the organic extract washed with water (until neutral) and brine, dried (MgSOA), filtered and concentrated to provide 58 mg of a solid.
- [9984] This material was acetylated (1.0 mL of dry pyridine, 0.20 mL of Ac<sub>2</sub>O, 23°C, N<sub>2</sub>, overnight), followed by workup (as described for the steroid of Example 14 flast step)) affording a crude product that was chromatographed (25 g SiO<sub>2</sub>, EtOAc). This product was triturated with ether to afford 51 mg (33%) of product as a white solid, m.p. 179-181°C.
  - [0085] MS (Cl. isobutane): (M+1) = 450 (M\*), 432, 391, 371, 348.
  - [0086] IR (KBr): 3398 (br), 2932, 2865, 1720 (sh. 1740), 1652, 1538, 1375, 1265, 1236 cm<sup>-1</sup>.
    - [0087] NMR (200 MHz 1H, CDCIs): 80.89, 1.22, 1.99, 2.17 (all s, 3H): 1.0-2.2 (m): 2.7 (t, J=13, 1H, H-8): 3.93 (s, 1H,
  - OH-17); 4.2 (br.s., 1H, H-11); 4.3 (br.s., 1H, H-3); 4.96 (AB, J=17.5, Av=42, 2H, H-21); 5.8 (d. J=10, 1H, NH).
  - [8088] Preferred embodiments of the present invention are the following embodiments denoted as Emb-1 to Emb-49:

# Emb-1. A compound of the following formula:

# Structure (A)

# Structure [8]

wherein R<sub>1</sub> is H, β-CH<sub>2</sub> or β-C<sub>2</sub>H<sub>5</sub>;

R<sub>2</sub> is F, C<sub>2</sub>-C<sub>11</sub> double bond, C<sub>2</sub>-C<sub>11</sub> epoxy, H or Cl;

 $F_0^1$  is  $I_1$ ,  $OR_{20}^{-}$ ,  $OC_1(=OR_{20}^{-})$ ,  $P_{20}^{-}$ , P

As is H, OH F, Cl, Br, CH<sub>3</sub>, phenyl, vinyl or allyl;

Re is H or CHa;

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Fig. Is CH<sub>2</sub>CH<sub>2</sub>OR<sub>26</sub>, CH<sub>2</sub>CH<sub>2</sub>OC(≈O)R<sub>27</sub>, H, OH, CH<sub>3</sub>, F, ≈CH<sub>2</sub>, CH<sub>2</sub>C(=O)OR<sub>29</sub>, OR<sub>26</sub>, 0(C=O)R<sub>27</sub> or O(C=O) CH<sub>2</sub>(C≈O)OR<sub>26</sub>

H<sub>10</sub> is -C∞CH, -CH-CH<sub>2</sub>, halogen, CN, N<sub>3</sub>, OR<sub>26</sub>, OC(=C)R<sub>27</sub>, H, OH, CH<sub>3</sub> or R<sub>10</sub> forms a second bond between positions C-16 and C-17;

R<sub>10</sub> is H or forms a double bond with R<sub>1</sub> or R<sub>14</sub>;

 $R_{13}$  is halogen,  $OR_{20}$ ,  $OC_1=OR_{27}$ ,  $NH_2$ ,  $NHG_{28}$ ,  $NHC(=OR_{27}$ ,  $N(R_{29})_2$ ,  $NC(=OR_{27}$ ,  $N_3$ , H. -OH, =O, -O-P(=O) (OH)2, or -O-C(=O)-(CH3)/COH where t is en integer from 2 to 6;  $R_{13}$  is H or forms a double bond with  $R_{13}$ :

R<sub>15</sub> Is H<sub>1</sub> =O or -OH;

and Rea with Rep forms a cyclic phosphate;

wherein R<sub>a</sub> and R<sub>18</sub> have the meaning defined above;

or wherein R<sub>23</sub> is -OH, O-C(=0)-R<sub>11</sub>, -OP(0)-(OH)<sub>2</sub>, or -O-C(=0)-(CH<sub>2</sub>)<sub>2</sub>COOH wherein t is an integer from 2 to 8; and R<sub>11</sub> is -Y-(CH<sub>2</sub>)<sub>n</sub>:X-(CH<sub>2</sub>)<sub>n</sub>:SO<sub>3</sub>H, -Y-(CH<sub>2</sub>)<sub>n</sub>:X-(CH<sub>2</sub>)<sub>n</sub>:NR<sub>16</sub>R<sub>17</sub> or -Z(CH<sub>2</sub>)<sub>n</sub>Q,

wherein Y is a bond or 0.5 Y is a bond, 0.5 or 0.5, and X is a bond, 0.5 Or 0.5, and 0.5 Or 0.

(1) -R<sub>19</sub>-CH<sub>2</sub>COOH wherein R<sub>19</sub> is -S-, -S(O)-, -S(O)<sub>2</sub>-, -SO<sub>2</sub>N(R<sub>20</sub>)-, or N(R<sub>20</sub>)SO<sub>2</sub>-; and R<sub>20</sub> is hydrogen or

lower alkyl-(C<sub>1</sub>-C<sub>4</sub>); with the proviso that the total number of carbon atoms in R<sub>20</sub> and (CH<sub>2</sub>), is not greater than 10: or

(2) -CO-COOH: or

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(3) CON(Ras)CH(Ras)COOH wherein Ras is H and Ras is H, CHa, -CHaCOOH, -CHaCHaCOOH, -CHaCOH, -CH<sub>2</sub>SH, -CH<sub>2</sub>CH<sub>2</sub>SCH<sub>2</sub>, or -CH<sub>2</sub>Ph-OH wherein Ph-OH is p-hydroxyphenyl;

or Ret is CH2 and Ree is H;

or Ro, and Roo taken together are -CHoCHoCHo-;

or -N(R<sub>21</sub>)CH(R<sub>22</sub>)COOH taken together is -NHCH<sub>2</sub>CONHCH<sub>3</sub>COOH; and pharmaceutically acceptable salts

with the proviso that except for the compound wherein R<sub>1</sub> is β-CH<sub>3</sub>, R<sub>2</sub> and R<sub>3</sub> taken together form a double bond between positions 9 and 11, R<sub>4</sub> and R<sub>6</sub> are hydrogen, R<sub>12</sub> and R<sub>14</sub> taken together form a double bond between positions 4 and 5, R<sub>5</sub> is α-F, R<sub>9</sub> is β-CH<sub>3</sub>, R<sub>10</sub> is α-OH, R<sub>13</sub> and R<sub>15</sub> are =O and R<sub>99</sub> is -OP(O)-(OH)<sub>2</sub>, R<sub>15</sub> is =O only when Rea with Ruo forms the above described cyclic phosphate.

 $R_{24} = C, C_1 - C_2$  double bond, O;

C(R15)CH2-R25, OH, OR26, OC(=0)R27, R26, COOH, C(=0)OR26, CHOHCH2OH, CHOHCH2OR26, CHOHCH, OC(=0)R27, CH, CH, CH, CH, CH, CH, CH, CH, CH, OC(=0)R27, CH, CN, CH, N3, CH, NH, CH2NHR26, CH2N(R26)2, CH2OH, CH2OR26, CH2O(C=O)R27, CH2O(P=O) (OH)2, CH2O(P=O) (OR26)2, CH2SH, CH2S-R26, CH2SC(=0)R27, CH2NC(=0)R27, C(=0)CHR26OH, C(=0)CHR26OR26, C(=0)  $CHR_{28}OC(=O)R_{27}$  or  $R_{10}$  and  $R_{28}$  taken together may be  $=C(R_{28})_2$ , that is, an optionally sixyl substituted methylene group;

wherein  $H_{26} = C_1 \cdot C_6$  (alkyl, branched alkyl, cycloalkyl, haloalkyl, aralkyl, aralkyl,  $H_{27} = H_{36} + OH_{26}$ ;  $H_{28} = H$ , C1-C8 25 (alkyl, branched alkyl, cycloalkyl); excepted from the compounds of Structure (A) are the compounds wherein R. is B-CH3 or B-C2H5;

Ro is H or Cl;

R<sub>3</sub> is H, =0, -OH, -O-alkyl(C<sub>1</sub>-C<sub>12</sub>), OC(=O)alkyl(C<sub>1</sub>-C<sub>12</sub>), -OC(=O)ARYL, -OC(=O)N(R)<sub>2</sub> or α-OC(=O)OR<sub>2</sub>, wherein ARYL is furyl, thienyl, pyrrolyl, or pyridyl and each of said moleties is optionally substituted with one or two (C,-C,)alkyl groups, or ARYL is -(CHo), phenyl wherein f is 0 to 2 and the phenyl ring is optionally substituted with t to 3 groups selected from chlorine, fluorine, bromine, alkyl(C<sub>4</sub>-C<sub>3</sub>), alkoxy(C<sub>4</sub>-C<sub>3</sub>), thicelkoxy-(C<sub>4</sub>-C<sub>3</sub>), Cl<sub>3</sub>C<sub>5</sub> F<sub>3</sub>C-, -NH<sub>2</sub> and -NHCOCH<sub>3</sub> and R is hydrogen, alkyl (C<sub>1</sub>-C<sub>4</sub>), or phenyl and each R can be the same or different, and R7 is ARYL as herein defined, or alkyl(C1-C12);

wherein R<sub>2</sub> and R<sub>3</sub> taken together are oxygen (-O-) bridging positions C-9 and C-11; or wherein R<sub>2</sub> and R<sub>3</sub> taken together form a double bond between positions C-9 and C-11; or Ro is α-F and Ro is β-OH;

or Ro is α-Cl and Ro is B-Cl;

and Ra Is H. CHa, Cl or F; 40 Rs Is H, OH, F, CI, Br, CHs, phenyl, vinyl or allyl;

Re is H or CHa:

Rolls H. OH, CHo. F or =CHo.

Rto is H, OH, CHa or Rto forms a second bond between positions C-16 and C-17;

R<sub>10</sub> is -H or forms a double bond with R<sub>14</sub>;

45 R<sub>13</sub> is H, -OH, =O, -O-P(O)(OH)<sub>2</sub>, or -O-C(=O)-(CH<sub>2</sub>)<sub>2</sub>COOH where t is an integer from 2 to 6;

R<sub>14</sub> is H or forms a double bond with R<sub>12</sub>;

R<sub>15</sub> is =0 or -OH;

and Res with Res forms a cyclic phosphate;

wherein Ro and Ros have the meaning defined above;

50 or wherein R2; is -OH O-C(=0)-R4, -OP(0)-(OH)2, or -O-C(=0)-(CH2)-COOH wherein t is an integer from 2 to 6; and R<sub>11</sub> is -Y-(CH<sub>2</sub>)<sub>0</sub>-X-(CH<sub>2</sub>)<sub>m</sub>-SO<sub>3</sub>H, -Y'-(CH<sub>2</sub>)<sub>0</sub>-X'-(CH<sub>2</sub>)<sub>0</sub>-NR<sub>16</sub>R<sub>17</sub> or -Z(CH<sub>2</sub>)<sub>1</sub>Q, wherein Y is a bond or -O-; Y' is a bond, -O-, or -S-; each of X and X' is a bond, -CON(R18)-, -N(R18)CO-, -O-, -S-, -S(O)-, or -S(O<sub>2</sub>)-, R18 is hydrogen or alkyl (C1-C4); each of R18 and R17 is a lower alkyl group of from 1 to 4 carbon atoms optionally substituted with one hydroxyl or R<sub>10</sub> and R<sub>12</sub> taken together with the nitrogen atom to which each is attached forms 55 a monocyclic heterocycle selected from pyrrolidino, piperidino, morpholino, thiomorpholino, piperazino or N(lower) alkyl-piperazino wherein alkyl has from 1 to 4 carbon atoms; n is an integer of from 4 to 9; m is an integer of from 1 to 5; p is an integer of from 2 to 9; q is an integer of from 1 to 5;

Z is a bond or -Q-; r is an integer of from 2 to 9; and Q is one of the following:

- (1) -R<sub>19</sub> CH<sub>2</sub>COOH wherein R<sub>18</sub> is -S<sub>7</sub> -S(O)<sub>7</sub>, -S(O)<sub>2</sub>, -SO<sub>2</sub>N(R<sub>20</sub>)-, or N(R<sub>20</sub>)SO<sub>2</sub>-; and R<sub>20</sub> is hydrogen or lower sikyl-(C<sub>1</sub>-C<sub>4</sub>); with the provise that the total number of carbon atoms in R<sub>20</sub> and (CH<sub>2</sub>)<sub>7</sub> is not greater than 10 or
- (2) -CO-COOH; or

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- (3) CON(R<sub>21</sub>)CH(R<sub>22</sub>)COOH wherein R<sub>21</sub> is H and R<sub>22</sub> is H, CH<sub>3</sub>, -CH<sub>2</sub>COOH, -CH<sub>2</sub>CH<sub>2</sub>COOH, -CH<sub>2</sub>OH, -CH<sub>2</sub>CH, -CH<sub>2</sub>CH<sub>2</sub>COOH, -CH<sub>2</sub>Ph-OH wherein Ph-OH is p-hydroxyphenyl;
  - or Roy is CH3 and Rog is H;
- or Ros and Ros taken together are -CHoCHoCHo-;
- 10 or N(R<sub>21</sub>)CH(R<sub>22</sub>)COOH taken together is -NHCH<sub>2</sub>CONHCH<sub>2</sub>COOH; and pharmaceutically acceptable salts theren!
  - with the proviso that except for the compound wherein  $R_1$  is  $\beta$ -CH $_3$ ,  $R_2$  and  $R_3$  taken together form a double bond between positions 9 and 11,  $R_1$  and  $R_2$  are hydrogen,  $R_12$  and  $R_{12}$  ands together form a double bond between positions 4 and 5,  $R_3$  is C-F,  $R_3$  is R-CH $_3$ ,  $R_1$ 0 is C-OH,  $R_1$ 3 and  $R_1$ 5 are = 0 and  $R_{12}$ 3 is = 0P(0)-(OH) $_2$ ,  $R_{13}$ 1 is = 00 only when  $R_{23}$ 3 with  $R_{13}$ 5 income the above described cyclic phosphate:
  - also excepted from the compounds of Siructure [A] are the compound 3.11 $\beta$ , 17 $\alpha$ , 21-terrahydroxy-5-pregnane-20-one (the 3- $\alpha$ , 5- $\beta$ , 3- $\alpha$ , 5- $\alpha$ , 3- $\beta$ , 5- $\alpha$ , and 3- $\beta$ , 5- $\beta$  is norm of tetrahydrocortical) wherein R<sub>19</sub> is -O. R<sub>10</sub> is  $\alpha$ -OH, R<sub>1</sub> is  $\beta$ -CH, R<sub>2</sub> is  $\beta$ -OH, R<sub>3</sub> is  $\beta$ -OH, Right observations, wherein R<sub>4</sub> is  $\beta$ -OH, R<sub>3</sub> is  $\beta$ -OH, Right of Signal (R<sub>3</sub>) is  $\beta$ -OH, R<sub>3</sub> is  $\beta$ -OH, Right of Signal (R<sub>3</sub>) is  $\beta$ -OH, R<sub>3</sub> is  $\beta$ -OH, Right of Signal (R<sub>3</sub>) is  $\beta$ -OH, Right of Signal (R<sub>3</sub>) is  $\beta$ -OH, R<sub>3</sub> is  $\beta$ -OH, Right of Signal (R<sub>3</sub>) is  $\beta$ -OH, R<sub>3</sub> is  $\beta$
  - dromostanolone propionate, wherein  $R_1$  is  $\beta$ -CH<sub>3</sub>,  $R_1R_1R_1R_2R_1R_3R_3R_3R_3$ , are  $H_1$ ,  $R_2$  is  $-\Theta$ .  $R_{12}$  is  $-\Theta$  and  $R_{22}$  is  $-\Theta$ .  $C(-\Theta)CH_2CH_2$ , methandrostenelone, wherein  $R_1$  is  $\beta$ -CH<sub>3</sub>,  $R_1$ - $R_1$ - $R_1$ - $R_2$ - $R_3$  is  $-\Theta$ . A wherein  $R_1$  is  $\beta$ -CH<sub>3</sub>,  $R_2$ - $R_3$ - $R_3$ - $R_3$ - $R_3$  is  $-\Theta$ . A collapse of  $R_2$  is  $P_1$ - $P_2$  is  $-P_2$ - $P_3$  is  $-P_2$ - $P_3$ - $P_3$
- and oxandrolone, wherein  $R_3$  is  $\beta$ -CH $_3$ ,  $R_2R_3R_5\bar{R}_6R_0R_{14}$  are H,  $R_{10}$  is  $\alpha$ CH $_3$ ,  $R_{12}$  is  $\alpha$ -H,  $R_{13}$  is =0,  $R_{24}$  is 0, and  $R_{26}$  is  $\beta$ -OH.
  - Emb-2. The compound of Emb- 1 selected from the group consisting of: 21-Nor-5β-pregnan-3α,17α,20-triol; 21-Nor-5β-pregnan-3α,17α,20-triol; 21-Nor-5β-pregnan-3α,17α,20-triol;
- 3-acetate: 21-Nor-Su-pregnan-Su, 17a, 20-triol-3-phosphate: 21-Nor-SB-pregn-17(20) en-3a, 18-triol: 21-Nor-SBpregnan-3a, 17B, 20-triol; 20-Acetamida-21-nor-SB-pregnan-3a, 17a-20-triol; 2-1-Nor-SB-pregnan-11β,17a,21-triol-20-one-21-acetate; 21-Nor-Sa-pregnan-3a, 17a,20-triol; 21a-Methyl-SB-pregnan-3a, 11β, 17a, 21-tatriol-20-one-21-methyl-ether; 20-Azido-21-nor-SB-pregnan-3a, 17a-diol; 2a(2-Ayetavy-pthyl-17β-methyl-sB-pregnan-3a, 17a-diol; 2a(2-Ayetavy-pthyl-17β-methyl-SB-pregnan-3a, 17a-diol; 2a(2-Ayetavy-pthyl-17β-methyl-SB-pregnan-3a, 17a-diol; 2a(2-Ayetavy-pthyl-17β-methyl-SB-pregnan-3a, 17a-diol; 2a(2-Ayetavy-pthyl-17β-methyl-SB-pregnan-3a, 17a-diol; 2a(2-Ayetavy-pthyl-17β-methyl-SB-pregnan-3a, 17a-diol; 2a(2-Ayetavy-pthyl-SB-pregnan-3a, 17a-diol; 2a(2
- 40 21-Nor-bj-pregn-1 /(20)en-3α-OL; 21-Nor-bj-pregn-1 /(20)en-3α-ol-3-acetate; 21-Nor-bj-pregn-1 /(20)-en-3α-ol-3-acetate; 3β-Azido-5β-pregnan-11β, 17α, 21-triol-20-one-21-acetate; and 5β-Pregnan-11β, 17α, 21-triol-20-one.
  - Emb- 3. The compound of Emb- 2 which is 21-Nor-5β-pregnan-3α,17α,20-triol and 21-Nor-5β-pregn-17(20)-en-3α,16-diol-3-acetate-16-(O-methyl) malonate.
    - Emb-4. A method for controlling ocular hypertension associated with primary open angle glaucoma which comprises administering a pharmaceutically effective amount of a compound of Emb-1.
- Emb-5. A method for controlling ocular hypertension associated with primary open angle glaucoma, which comprises administering a pharmaceutically effective amount of a compound selected from the group consisting of: 21-Nor-5β-pregnan-3c,17c,20-triol-3-phosphate; 21-Nor-5β-pregna-7c,17c,20-triol-3-phosphate; 21-Nor-5β-pregna-7c,17c,20-triol-3-phosphate; 21-Nor-5β-pregna-3c,17c,20-triol-3-phosphate; 21-Nor-5β-pregna-3c,17c,20-triol-3-phosphate; 21-Nor-5β-pregna-3c,17c,20-triol-3-phosphate; 21-Nor-5β-pregna-3c,17c,20-triol-3-phosphate; 21-Nor-5β-pregna-3c,17c,20-triol-3-phosphate; 21-Nor-5β-pregna-3c,17c,3c-triol-3c-phosphate; 21-Nor-5β-pregna-3c,17c,3c-triol-3c-phosphate; 21-Nor-5β-pregna-3c,17c-diol-3-phosphate; 21-Nor-5β
- c): 20(Carbethoxymethyl)thio-21-nor-5β-pregnan-3α,17α-diol; 20-(4-Fluorophenyl)thio-21-nor-5β-pregnan-3α, 17α-diol; 16α-(2-Hydroxyethyl)-1/β-methyl-5β-androsian-3α,17α-diol; 20-Cyano-21-nor-5β-pregnan-3α,17α-diol: 17α-Methyl-5β-androsian-3α,17α-diol: 17α-diol: 1

3-acetate; 21-Nor-5β-pregn-17(20)-en-3α-ol-16-acetic acid 3-acetate; 3β-Azido-5β-pregnan-11β, 17α, 21-triol-20-one-21-acetate; and5β-pregnan-11β, 17α, 21-triol-20-one-4,9(11)-Pregnadien-17α,21-diol-3\_20-dione 21-acetate; 4-Androstan-3-one-17β-carboxylic acid; 17α-Ethynyl-51(1)-acten-17β-ol-3-one; 17α-Ethynyl-1,3,5(10)-actratrien-3,17β-diol; 11-Epicortisol; 17α-Hydroxyprogesterone; and Tetrahydrocortoxyline.

Emb-6. The method of Emb-5 wherein the compound is 4, 9(11)-Pregnadien-17a, 21-diol-3, 20-dione-21-acetate.

Emb-7. A composition for controlling ocular hypertension associated with primary open angle glaucoma comprising a pharmaceutically effective amount of a compound of Emb-1.

Emb. 3. A composition for controlling ocular hypertension comprising a pharmaceutically effective amount of a compound scienced from the group consisting of: 21-Nor-5β-pregnan-3α, 17α,20-triol-3-breathe; 21-Nor-5β-pregnan-3α, 17α,20-triol-3-breathe; 21-Nor-5β-pregnan-17(2)0-no-2, 17α,20-triol-3-breathe; 21-Nor-5β-pregnan-3α, 17β,20-triol; 20-dectamide 21-nor-5β-pregnan-3α, 17β,20-triol; 21-delleh)-3-deptengran-11,17,17α,21-triol-20-one-21-activity ather; 20-Azido 21-nor-5β-pregnan-3α, 17α-diol; 21-delleh)-17α,17α,21-triol-20-one-21-activity ather; 20-Azido 21-nor-5β-pregnan-3α, 17α-diol; 21-delleh)-17α-diol; 21-nor-5β-pregnan-3α, 17α-diol; 20-de-Elucrophylihio-21-nor-5β-pregnan-3α, 17α-diol; 20-de-Elucrophylihio-21-nor-5β-pregnan-3α, 17α-diol; 17α-Methyl-5β-androstan-3α, 17α-diol; 21-Nor-5β-pregnan-11,17α,01-α-diol; 21-Nor-5β-pregnan-11,17α,01-α

Emb-9. The composition of Emb-7 wherein the compound is present at a concentration between 0.005 and 5.0 weight percent.

Emb-10. The composition of Emb-8 wherein the compound is 4, 9(11)-Pregnadien-17α,21-diol-3,20-dione-21-acetate,

Emb-11. The composition of Emb- 8 wherein the compound is present at a concentration of between 0.005 and 5.0 weight percent.

26 Emb-12. A composition for preventing and treating neovascularization comprising a therapeutically effective amount of an angiostatic steroid of the following formula:

Structure [A] Structure [B]

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each of said moleties is optionally substituted with one or two  $(C_1, C_2)$ alityl groups, or ARYL is  $-(CH_2)$ -phenyl wherein f is 0 to 2 and the phenyl ring is optionally substituted with 1 to 3 groups selected from chlorine, fluorine, bromine, alisyl( $C_1, C_2$ ), allkoy( $C_1, C_2$ ), thiosalkoy+ $(C_1, C_2)$ ,  $C_2$ ,  $C_2$ ,  $C_3$ ,  $C_3$ ,  $C_4$ , C

R<sub>8</sub> is H, OH, F, Cl, Br, CH<sub>9</sub>, phenyl, vinyl or allyl;

Rs is H or CH3;

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 $\begin{array}{l} R_{9} \text{ is } CH_{2}CH_{2}OR_{28}, CH_{2}CH_{2}OC(=O)R_{27}, H, OH, CH_{3}, F, =CH_{2}, \text{ or } CH_{2}C(=O)OR_{26}, OR_{26}, O(C=O)R_{27}, \text{ or } O(C=O)CH_{2}(C=O)OR_{26}, OR_{26}, OR_$ 

R<sub>10</sub> is -C=CH, -CH=CH<sub>2</sub>, halogen, CN, N<sub>3</sub>, OR<sub>28</sub>, OC(=O)R<sub>27</sub>, H, OH, CH<sub>3</sub> or R<sub>10</sub> forms a second bond between positions C-18 and C-17;

R<sub>12</sub> is H or forms a double bond with R<sub>1</sub> or R<sub>14</sub>;

 $R_{13} \text{ is halogen, } OR_{26}, OC(=0)R_{27}, NH_{2}, NHR_{26}, NHC(=0)R_{27}, N(R_{26})_2, NC(=0)R_{27}, N_{3}, H, -OH, =O, -O-P(=O) + (OH)_{2}, or -O-C(=O)-(CH_{2})(COOH) where t is an integer from 2 to 6;$ 

R<sub>14</sub> is H or forms a double bond with R<sub>12</sub>;

R<sub>15</sub> is H, =0 or -0H;

and R23 with R10 forms a cyclic phosphate;

wherein Rs and Rs have the meaning defined above;

or wherein R<sub>23</sub> is -OH, O-C(=O)-R<sub>11</sub>, -OP(O)-(OH)<sub>2</sub>, or -O-C(=O)-(CH<sub>2</sub>)<sub>1</sub>COOH

20 wherein t is an integer from 2 to 6; and R<sub>11</sub> is -Y-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>m</sub>-SO<sub>3</sub>H, -Y'(CH<sub>2</sub>)<sub>p</sub>-X'-(CH<sub>2</sub>)<sub>q</sub>-NR<sub>1g</sub>R<sub>17</sub> or -Z (CH<sub>2</sub>)<sub>p</sub>Q.

wherein Y is a bond or -0: Y is a bond. -0, or -8; each of X and X is a bond. -0CN( $R_{18}$ ),  $-N(R_{18})$ CC-, -0. S. -8(O)-, or -8(O)-, -1R<sub>1</sub> is hydrogen or alky! (C<sub>1</sub>-C<sub>2</sub>), each of  $R_{18}$  is clower alify group of from 1 to 4 carbon atoms potitionally substituted with one hydroyl or  $R_{18}$  and  $R_{17}$  take not opether with the nitrogen atom to which each is attached forms a monocyclic heterocycle selected from pyrrolldino, piperdino, morpholino, thiomorpholino, piperazino or N(lower)alkyl-piperazino wherein alkyl has from 1 to 4 carbon atoms; n is an integer of from 4 to 8; m is an integer of from 2 to 8; is an integer of from 3 to 8; is an integer of from 5 to 8; in the proper of the first of 8 is an integer of from 2 to 8.

Z is a bond or -O-; r is an integer of from 2 to 9; and Q is one of the following:

 -R<sub>19</sub>:CH<sub>2</sub>COOH wherein R<sub>19</sub> is -S-, -S(O)-, -S(O)<sub>2</sub>-, -SO<sub>2</sub>N(R<sub>20</sub>)-, or N(R<sub>20</sub>)SO<sub>2</sub>-; and R<sub>20</sub> is hydrogen or lower alkyl-(C<sub>1</sub>-C<sub>4</sub>); with the provise that the total number of carbon atoms in R<sub>20</sub> and (CH<sub>2</sub>), is not greater than 10: or

(2) -CO-COOH: or

(3) CON(R<sub>21</sub>)CH(R<sub>22</sub>)COOH wherein R<sub>21</sub> is H and R<sub>22</sub> is H, CH<sub>3</sub>, -CH<sub>2</sub>COOH, -CH<sub>2</sub>CH<sub>2</sub>COOH, -CH<sub>2</sub>OH, -CH<sub>2</sub>CH<sub>3</sub>CH<sub>4</sub>CH<sub>5</sub>CH<sub>6</sub>, or -CH<sub>2</sub>Ph-OH wherein Ph-OH is p-hydroxyphenyl,

or R21 is CH3 and R22 is H;

or R21 and R22 taken together are -CH2CH2CH2-:

or -N(R<sub>21</sub>)CH(R<sub>22</sub>)COOH taken together is -NHCH<sub>2</sub>CONHCH<sub>2</sub>COOH; and pharmaceutically acceptable selts thereof:

with the proviso that except for the compound wherein  $R_1$  is  $-CH_3$ ,  $R_2$  and  $R_3$  taken together form a double bond between positions 9 and 11,  $R_2$  and  $R_3$  are hydrogen,  $R_{12}$  and  $R_{14}$  laken together form a double bond between positions 4 and 5,  $R_3$  is  $-F_1$ ,  $R_3$  is  $-CH_3$ ,  $R_{10}$  is  $-CH_3$ ,  $R_{13}$  and  $R_{13}$  are -C and  $R_{23}$  is -CP(O)- $(CH)_2$ ,  $R_{13}$  is -C only when  $R_{23}$  with  $R_{13}$  forms the above described cyclic phosphate;

 $R_{24} = C, C_1 - C_2$  double bond, O;

 $R_{29} = (R_{13}^{\circ}Gl_{24}^{\circ}R_{29}, OH, OR_{20}, OC(=O)R_{27}, R_{20}, COOH, C(=O)R_{29}, CHOHCH_2OH, CHOHCH_2OR_{20}, CHOHCH_2OC(=O)R_{27}, CH_2OH, OH, CH_2OH_2OR_{20}, CH_2CH_2OC(=O)R_{27}, CH_2OH, OH_2OH, CH_2OH, C$ 

R<sub>25</sub> taken together may be «C(R<sub>26</sub>)<sub>2</sub>, that is, an optionally alkyl substituted methylene group;

wherein  $H_{26} = C_1 - C_6$  (alkyl, branched alkyl, cycloalkyl, haloalkyl, aralkyl, aryl);  $H_{27} = H_{26} + OR_{26}$ ;  $H_{26} = H$ . C1-C8 (alkyl, branched alkyl, cycloalkyl); excepted from the compounds of Structure [A] are the compounds wherein  $H_1$  is F-CH<sub>4</sub> or F-CH<sub>4</sub>.

Rols Hor -CI;

 $R_3 \text{ is =0, -OH, -O-alkyl}(C_1-C_{12}), -OC(=0) \text{alkyl}(C_1-C_{12}), -OC(=0) \text{ARYL, -OC}(=0) \text{N(R)}_2 \text{ or } \alpha -OC(=0) \text{OR}_2, \text{ wherein } \alpha \in \mathbb{R}^n$ 

ARYL is turyl, thienyl, pyrrolyl, or pyridyl and each of said moietles is optionally substituted with one or two (C<sub>1</sub>-C<sub>s</sub>) alkyl groups, or ARYL is -(CH<sub>2</sub>), phenyl wherein f is 0 to 2 and the phenyl ring is optionally substituted with 1 to 3 groups selected from chilorine, fluorine, bromine, alkyl(C<sub>1</sub>-C<sub>2</sub>), alkoxy(C<sub>4</sub>-C<sub>3</sub>), thioalkoxy-(C<sub>1</sub>-C<sub>3</sub>), Cl<sub>3</sub>C-, F<sub>3</sub>C-, -NH<sub>2</sub> and -NHCOCH, and R is hydrogen, alkyl (C,-C,4), or phenyl and each R can be the same or different, and R, is ARYL as herein defined, or alkyl(C<sub>4</sub>-C<sub>49</sub>):

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wherein R<sub>2</sub> and R<sub>3</sub> taken together are oxygen (-O-) bridging positions C-9 and C-11; or wherein B<sub>2</sub> and B<sub>3</sub> taken together form a double bond between positions C-9 and C-11: or Ro is ex-F and Ro is B-OH;

or Ro is a Cl and Ro is B-Cl;

and R<sub>4</sub> is H. CH<sub>5</sub>, Cl or F:

Rs is H, OH, F, Ci, Br, CH3, phenyl, vinyl or allyl;

Re is H or CHa;

Ra is H, OH, CH3, F or =CH5;

R<sub>10</sub> is H, OH, CH<sub>2</sub> or R<sub>10</sub> forms a second bond between positions C-16 and C-17;

R<sub>12</sub> is -H or forms a double bond with R<sub>14</sub>;

R<sub>13</sub> is H, -OH, =O, -O-P(O)(OH)<sub>2</sub>, or -O-C(=O)-(CH<sub>2</sub>),COOH where t is an integer from 2 to 6;

R14 is H or forms a double bond with R19; R<sub>15</sub> is =0 or -OH;

and Rea with Rin forms a cyclic phosphate;

wherein Re and Res have the meaning defined above;

or wherein Res is -OH, O-C(=O)-Res, -OP(O)-(OH)-, or -Q-C(=O)-(CH-),COOH

wherein t is an integer from 2 to 6; and R<sub>11</sub> is -Y-(CH<sub>2</sub>)<sub>a</sub>-X-(CH<sub>2</sub>)<sub>a</sub>-SO<sub>3</sub>H, -Y'(CH<sub>2</sub>)<sub>a</sub>-X'-(CH<sub>2</sub>)<sub>a</sub>-NR<sub>16</sub>R<sub>17</sub> or -Z (CHo), Q, wherein Y is a bond or -O-; Y' is a bond, -O-, or -S-; each of X and X' is a bond, -CON(R, a)-, -N(R, a)CO-. -O-, -S-, -S(O)-, or -S(O<sub>2</sub>)-; R<sub>18</sub> is hydrogen or alkyl (C<sub>1</sub>·C<sub>4</sub>); each of R<sub>16</sub> and R<sub>17</sub> is a lower alkyl group of from 1 to 4 carbon atoms optionally substituted with one hydroxyl or Ris and Riz taken together with the nitrogen atom

to which each is attached forms a monocyclic heterocycle selected from pyrrolidino, piperidino, morpholino, thiomorpholino, piperazino or N(lower)alkyl-piperazino wherein alkyl has from 1 to 4 carbon atoms; n is an integer of from 4 to 9; m is an integer of from 1 to 5; p is an integer of from 2 to 9; q is an integer of from 1 to 5; Z is a bond or -O-; r is an integer of from 2 to 9; and Q is one of the following:

- -R<sub>19</sub>-CH<sub>2</sub>COOH wherein R<sub>19</sub> is -S-, -S(O)-, -S(O)<sub>2</sub>-, -SO<sub>2</sub>N(R<sub>20</sub>)-, or N(R<sub>20</sub>)SO<sub>2</sub>-; and R<sub>20</sub> is hydrogen or lower alkyl-(C1-C4); with the provise that the total number of carbon atoms in R20 and (CH2), is not greater than 10; or
- (2) -CO-COOH: pr
- (9) CON(R<sub>21</sub>)CH(R<sub>22</sub>)COOH wherein R<sub>21</sub> is H and R<sub>22</sub> is H, CH<sub>31</sub> -CH<sub>3</sub>COOH, -CH<sub>2</sub>COOH, -CH<sub>2</sub>COOH, -CH<sub>2</sub>OOH, -CHoSH, -CHoCHoSCHo, or -CHoPh-OH wherein Ph-OH is p-hydroxyphenyl;
- or Ray is CHa and Ray is H;
- or R21 and R22 taken together are -CH2CH2CH2-;
  - or -N(R<sub>21</sub>)CH(R<sub>20</sub>)COOH taken together is -NHCH<sub>2</sub>CONHCH<sub>3</sub>COOH; and pharmaceutically acceptable salts

with the provise that except for the compound wherein R, is \$-CHa, Re and Ra taken together form a double bond between positions 9 and 11, Re and Re are hydrogen, R12 and R14 taken together form a double bond between positions 4 and 5, R<sub>5</sub> is α-F, R<sub>9</sub> is β-CH<sub>9</sub>, R<sub>10</sub> is α-OH, R<sub>13</sub> and R<sub>15</sub> are =0 and R<sub>23</sub> is -OP(0)-(OH)<sub>2</sub>, R<sub>13</sub> is =0 only when R23 with R10 forms the above described cyclic phosphate;

also excepted from the compounds of Structure [A] are the compounds 3,11 ß, 17a, 21-tetrahydroxy-5 -pregnane-20-one (the 3-alpha, 5-beta; 3-alpha, 5-alpha; 3-beta, 5-alpha; and 3-beta, 5-beta isomers of tetrahydrocortisol) wherein R<sub>1x</sub> is =0, R<sub>10</sub> is α OH, R<sub>1</sub> is CH<sub>2</sub>, R<sub>2</sub> is β OH, R<sub>2</sub> is H, R<sub>4</sub> is H, R<sub>12</sub> is α or β OH, R<sub>14</sub> is H, R<sub>10</sub> is α or β

50 H, Re is H, Re is H, Ro is H, Rot is C, and R23 is OH.

> Emb-13. A composition for preventing or treating neovascularization, comprising: a therapeutically effective amount of an anglostatic steroid selected from the group consisting of: 21-Nor-5\$-pregnan-3a,17a,20-triol-3-scetate; 21-Nor-5α-pregnan-3α,17α,20-triol-3-phosphate; 21-Nor-5β-pregn-17(20)en-3α,16-diol;21-Nor-5β-pregnan-3a,178,20-triol;20-Acetamide-21-nor-58-pregnan-3a,17a-diol-3-acetate; 38 Acetamido-58-pregnan-118, 17α,21-triol-20-one-21-acetate;21-Nor-5α-pregnan-3α,178.20-triol;21α-Methyl-5β-pregnan-3α,118, 17α, 21-tetrol-20-one-21-methyl ether; 20-Azido-21-nor-58-pregnan-3α, 17α-diol; 20(Carbethoxymethyl)thio-21-nor-58-pregnan-3α.17α-diol. 20-(4-Fluorophenyl)thio-21-nor-56-pregnan-3α.17α-diol. 16α-(2-Hydroxyethyl)-178-methyl-56-

androstan-Sq. 17 $\alpha$ -diol; 20-Cyano-21-nor-5 $\beta$ -pregnan-Sq. 17 $\alpha$ -diol; 17 $\alpha$ -Methyl-5 $\beta$ -androstan-Sq. 17 $\beta$ -diol; 21-Nor-5 $\beta$ -pregn-17(20)en-S $\alpha$ -Ol.; 21-Nor-5 $\alpha$ -ol-sacetate; 21-Nor-5 $\alpha$ -ol-sacetate; 3 $\beta$ -Azido-5 $\beta$ -pregnan-11 $\beta$ , 17 $\alpha$ . 21-triol-20-one-21-acetate; and 5 $\beta$ -pregnan-11 $\beta$ , 17 $\alpha$ . 21-triol-20-one-4, 9(11)-Pregnedien-17 $\alpha$ , 21-diol-3, 20-dione; 4, 9(11)-Pregnedien-17 $\alpha$ , 21-diol-3, 20-dione; 4, 9(11)-Pregnedien-17 $\alpha$ , 21-diol-3, 20-dione; 4, 9(11)-Pregnedien-17 $\alpha$ , 21-diol-3, 20-dione; 4)-diol-3, 21-diol-3, 21-diol-3,

- Emb-14. The composition of Emb-12 wherein the anglostatic steroid concentration is 0.01 10.0 wt.%.
- 10 Emb-15. The composition of Emb-14 wherein the concentration is 0.1 5.0 wt.%.
  - Emb-18. A method for preventing and treating neovascularization, which comprises: administering a therapeutically effective amount of the composition of Emb-12.
- 16 Emb-17. The method for preventing and treating neovascularization which comprises: administering a therapeutically effective amount of the composition of Emb-13.
  - Emb-18. The method of Emb-16 wherein the neovascularization being prevented and treated is selected from the group consisting of head traums, along interest and treated in selected from the group consisting of head traums, and interest and interest and the selected from the group consisting of the selected from the group cannot be along a factor of the selected from the group cannot be along the selected from the group cannot be group cannot be along the selected from the group cannot be g
- 25 Emb-19. A method for preventing and treating ocular neovascularization, which comprises: administering a therapeutically effective amount of the composition of Emb-12.
  - Emb-20. A method for preventing and treating ocular neovascularization, which comprises: administering a therapeutically effective amount of the composition of Emb-13.
  - Emb-21. The method of Emb- 19 wherein the angiostatic steroid concentration is 0.01 10 wt.%.
    - Emb-22. The method of Emb-19 wherein the angiostatic steroid concentration is 0.1 to 5.0 wt.%.
- 35 Emb-23. The method of Emb-19 wherein the ocular neovascularization being prevented and treated is in the front of the eve.
  - Emb-24. The method of Emb-19 wherein the ocular neovascularization being treated is in the cornea.
- 40 Emb- 25. The method of Emb-19 wherein the ocular neovascularization being prevented and treated is in the back of the eye.
  - Emb- 26. The method of Emb-17 wherein the angiostatic steroid is 4,9(11)-Pregnadien-17a,21-diol-3,20-dione-21-acetate.
  - Emb-27. The method of Emb-17 wherein the anglostatic steroid is 4,9(11)-Pregnadien-17a,21-diol-3,20-dione.
    - Emb-28. The method of Emb-17 wherein the angiostatic steroid is administered at a concentration of about 0.01 to 10.0 weight percent.
    - Emb-29. The method of Emb-17 wherein the angiostatic steroid is administered at a concentration of about 0.1 5.0 weight percent.
- Emb-30. A method for preventing and treating neovascularization of the tissues in the front of the eye, which comprises; administering a pharmaceutically effective amount of a composition of Emb-13.
  - Emb-31. The method of Emb-30 wherein the angiostatic steroid is 4,9(11)-Pregnadien-17a,21-diol-3,20-dione-21-acetats.

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Emb-32. The method of Emb- 30 wherein the angiostatic steroids is 4,9(11)-Pregnadien-17a,21-diol-3,20-dione.

Emb-33. The method of Emb-30 wherein the angiostatic steroid is administered at a concentration of about 0.01 to 10.0 weight percent.

Emb-34. The method of Emb-31 wherein the concentration is about 0.1 - 5.0 weight percent.

Emb-35. The method of Emb-30 wherein the tissue being treated is the comea.

6 Emb-36. A method for preventing and treating neovascularization of the tissues of the back of the eye, which comprises; administering a pharmaceutically effective amount of a composition of Emb-13.

Emb-37. The method of Emb-36 wherein the angiostatic steroid is 4.9(11)-Pregnadien-17a,21-diol-3.20-dione.

Emb-38. The method of Emb-36 wherein the angiostatic steroid is administered at a concentration of about 0.01 to 10.0 weight percent.

Emb-39. The method of Emb-38 wherein the angiostatic sterold is administered at a concentration of about 0.1 to 5.0 weight percent.

Emb-40. The method of Emb-36 wherein the tissue being treated is the retina or the subretina.

Emb-41. The method of Emb-36 wherein the tissue being treated is the macula.

25 Emb-42. The method of Emb-36 wherein the tissue being treated is the optic nerve head.

Emb-43. A pharmaceutical composition useful in the treatment of ophthalmic inflammation, comprising: an antiinflammatiory effective amount of a glucocorticoid and an intraocular pressure controlling amount of an anglostatic steroid of the formula:

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Structure (A)

Structure [B]

wherein R<sub>4</sub> is H<sub>1</sub> 8-CH<sub>2</sub> or B-C<sub>2</sub>H<sub>6</sub>;

Rg is F, Cg-C11 double band, Cg-C11 epaxy, H or Cl;

 $R_0$  is  $H_1$ ,  $GR_{g_0}$ :  $O(c(-G)R_{2T_1}$ , halogen,  $C_g - C_{11}$  double bond,  $C_g - C_{11}$ , epoxy, = O,  $-OH_1$ ,  $(-O-B)K/(C_1 - C_2)$ ,  $>O(c(-O)BRV_1)$  and each of said moleties is optionally substituted with one or two  $(C_1 - C_2)aKy$  groups, or ARYL is  $-(CH_2)aKy$  therein A is  $-(CH_2)aKy$  groups, or ARYL is  $-(CH_2)aKy$  therein A is  $-(CH_2)aKy$  groups, and  $-(C_1 - C_2)aKy$  groups selected from chlorine, fluorine, bromine,  $-(C_1 - C_2)aKy$  ( $-(C_1 - C_2)aKy$ ) and  $-(C_1 - C_2)aKy$  and  $-(C_1 - C_2)AK$ 

Rs is H. OH, F, Cl, Br, CH<sub>3</sub>, phenyl, vinyl or allyl;

Re is H or CHs:

R<sub>q</sub> is CH<sub>2</sub>CH<sub>2</sub>OR<sub>20</sub>, CH<sub>3</sub>CH<sub>5</sub>OC(≈O)R<sub>27</sub>, H, OH, CH<sub>9</sub>, F, ≈CH<sub>6</sub>, CH<sub>2</sub>C(≈O)OR<sub>28</sub>, OR<sub>28</sub>, O(C≈O)R<sub>37</sub> or O(C=O)

CH2(C=O)OR26

R<sub>10</sub> is -C=-CH<sub>2</sub>-CH=CH<sub>2</sub>, halogen, CN, N<sub>3</sub>, OR<sub>26</sub>, OC(=O)R<sub>27</sub>, H, OH, CH<sub>3</sub> or R<sub>10</sub> forms a second bond between positions C-18 and C-17;

R<sub>12</sub> is H or forms a double bond with R<sub>1</sub> or R<sub>14</sub>;

5 R<sub>13</sub> Is halogen, OR<sub>26</sub>, OC(=0)R<sub>27</sub>, NH<sub>2</sub>, NHR<sub>26</sub>, NHC(=0)R<sub>27</sub>, N(R<sub>26</sub>)<sub>2</sub>, NC(=0)R<sub>27</sub>, N<sub>3</sub>, H, -OH, =O, -O-P(=0) (OH)<sub>2</sub>, or -O-C(=0)-(CH<sub>2</sub>)<sub>2</sub>COOH where t is an integer from 2 to 6;

Rea is H or forms a double bond with Rea;

R<sub>15</sub> is H, = O or -OH;

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and R<sub>23</sub> with R<sub>10</sub> forms a cyclic phosphate;

wherein  $R_9$  and  $R_{15}$  have the meaning defined above; or wherein  $R_{23}$  is -OH, O-C(=O)- $R_{11}$ , -OP(O)-(OH)<sub>2</sub>, or -O-C (=O)-(CH<sub>3</sub>),COOH

wherein t is an integer from 2 to 6; and  $R_{11}$  is  $-Y-(CH_2)_n$   $X-(CH_2)_m$   $SO_3H$ ,  $-Y-(CH_2)_p$   $X-(CH_2)_q$   $NR_{16}R_{17}$  or -Z  $(CH_2)_q$ .

whorein Y is a bond or -0- Y is a bond, -0-, or -8-, each of X and X is a bond,  $-CON(R_{18})$ ,  $-N(R_{18})CO$ -, -C-, -8-, -

Z is a bond or -O-; r is an integer of from 2 to 9; and Q is one of the following:

(1)  $-R_{19}$ : CH<sub>2</sub>COOH wherein  $R_{19}$  is  $-S_1$ ,  $-S(O)_2$ ,  $-S(O)_2$ ,  $-S_2N(R_{20})_7$ , or  $N(R_{20})SO_2$ , and  $R_{20}$  is hydrogen or lower alloyI-(C<sub>1</sub>-C<sub>4</sub>), with the proviso that the total number of carbon atoms in  $R_{20}$  and (CH<sub>2</sub>), is not greater than 10; or

(2) -CO-COOH: or

(3) CON(R<sub>21</sub>)CH(R<sub>22</sub>)COOH wherein R<sub>21</sub> is H and R<sub>22</sub> is H, CH<sub>3</sub>. -CH<sub>2</sub>COOH, -CH<sub>2</sub>CH<sub>2</sub>COOH, -CH<sub>2</sub>OH, -CH<sub>2</sub>CH<sub>2</sub>COOH, -CH<sub>2</sub>CH<sub>2</sub>COOH, -CH<sub>2</sub>OH, -CH<sub>2</sub>CH<sub>2</sub>COOH, -CH<sub>2</sub>COOH, -CH<sub></sub>

or R21 is CH3 and R22 is H;

30 or R<sub>21</sub> and R<sub>22</sub> taken together are -CH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>·· or -N(R<sub>21</sub>)CH(R<sub>22</sub>)COOH taken together is -NHCH<sub>2</sub>COHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHCH<sub>2</sub>COOHC

R<sub>24</sub> = C, C<sub>1</sub>-C<sub>2</sub> double bond, O;

 $\begin{array}{lll} \overrightarrow{R_{25}} = & \overrightarrow{C(R_{19})}\overrightarrow{CH_{2}}.R_{23}, & OH, & \overrightarrow{CR_{29}}, & OC(=O)R_{27}, & R_{28}, & OCOH, & O(=O)OR_{29}, & OHOHCH-JOH, & CHOHCH-JOR_{29}, \\ & CHOHCH-JOC(=O)R_{27}, & CH_2JOH, & CH_2JOH_{29}, & CH_2CH_2OC(=O)R_{27}, & CH_2CH, & CH_2JOH, \\ & CH_3JOHR_{29}, & CH_4JOHR_{39}, & CH_4JOH, & CH_{39}, & CH_4JOHR_{39}, & CH_4JOHR_{39},$ 

49 wherein R<sub>20</sub> = C<sub>1</sub>-C<sub>2</sub> (alityl, tranched alkyl, cycloalityl, haloalityl, arallyl, aryl); R<sub>27</sub> = R<sub>26</sub> + OR<sub>26</sub>, R<sub>26</sub> = H, C1-C8 (alityl, branched alkyl, cycloalityl); excepted from the compounds of Structure (A) are the compounds wherein R<sub>1</sub> is β-CH<sub>2</sub> or β-C<sub>2</sub>H<sub>3</sub>;
R<sub>3</sub> is H or C1;

55 wherein R<sub>2</sub> and R<sub>3</sub> taken together are oxygen (O-) bridging positions C-9 and C-11; or wherein R<sub>2</sub> and R<sub>3</sub> taken together form a double bond between positions C-9 and C-11; or R<sub>3</sub> is x-F and R<sub>4</sub> is B-OH;

or R2 is a-CI and R3 is \$-CI;

and Ra is H, CH3, Cl or F; Rs is H, OH, F, Cl. Br, CH<sub>3</sub>, phenyl, vinyl or allyl; Re is H or CHa:

Ro is H. OH. CHa, F or = CHa;

5 Rue is H. OH, CHe or Rue forms a second bond between positions C-16 and C-17;

R12 is -H or forms a double bond with R14:

R<sub>10</sub> is H<sub>1</sub> -OH<sub>2</sub> = O<sub>1</sub> -O-P(O)(OH)<sub>0</sub>, or -O-C(=O)-(CH<sub>0</sub>),COOH where t is an integer from 2 to 6;

Ris H or forms a double bond with Ris

Ats is = O or -OH;

10 and Ros with Ros forms a cyclic phosphate;

wherein R<sub>9</sub> and R<sub>15</sub> have the meaning defined above; or wherein R<sub>23</sub> is -OH, O-C(=O)-R<sub>11</sub>, -OP(O)-(OH)<sub>21</sub> or -O-C (=0)-(CH<sub>2</sub>),COOH

wherein t is an integer from 2 to 6; and R<sub>11</sub> is -Y-(CH<sub>2</sub>)<sub>n</sub>-X-(CH<sub>2</sub>)<sub>m</sub>-SO<sub>3</sub>H, -Y'-(CH<sub>2</sub>)<sub>e</sub>-X'-(CH<sub>2</sub>)<sub>0</sub>-NR<sub>16</sub>R<sub>17</sub> or -Z (CH<sub>2</sub>),Q, wherein Y is a bond or Q; Y is a bond, Q, or S; each of X and X is a bond, CON(R<sub>18</sub>), N(R<sub>18</sub>)CO. 15 -O-, -S-, -S(O)-, or -S(O<sub>2</sub>)-; R<sub>18</sub> is hydrogen or alkyl (C<sub>1</sub>-C<sub>4</sub>)] each of R<sub>18</sub> and R<sub>17</sub> is a lower alkyl group of from 1 to 4 carbon atoms optionally substituted with one hydroxyl or R<sub>16</sub> and R<sub>17</sub> taken together with the nitrogen atom to which each is attached forms a monocyclic heterocycle selected from pyrrolldino, piperidino, morpholino, thismorpholino, piperazino or N(lower)alkyl-piperazino wherein alkyl has from 1 to 4 carbon atoms; n is an integer of from 4 to 9; m is an integer of from 1 to 5; p is an integer of from 2 to 9; q is an integer of from 1 to 5; 20

Z is a bond or -O-; r is an integer of from 2 to 9; and Q is one of the following:

 -R<sub>19</sub>-CH<sub>2</sub>COOH wherein R<sub>19</sub> is -S<sub>7</sub>, -S(O)<sub>7</sub>, -S(O)<sub>2</sub>, -SO<sub>2</sub>N(R<sub>20</sub>)-, or N(R<sub>20</sub>)SO<sub>2</sub>, and R<sub>20</sub> is hydrogen or lower alkyl-(C<sub>1</sub>-C<sub>2</sub>); with the proviso that the total number of carbon atoms in R<sub>20</sub> and (CH<sub>2</sub>), is not greater than 10; or

(2) -CO-COOH: or

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(3) CON(Reg)CH(Reg)COOH wherein Reg is H and Reg is H, CH<sub>3</sub>, -CH<sub>2</sub>COOH, -CH<sub>2</sub>COOH, -CH<sub>2</sub>COOH, -CH<sub>2</sub>OOH, -CH2SH, -CH2CH2SCH3, or -CH2Ph-OH wherein Ph-OH is p-hydroxyphenyl;

or Ro, is CHa and Roals H;

or R21 and R22 taken together are -CH2CH2CH2-;

or -N/Ras/CH/Ras/COOH taken together is -NHCH-CONHCH-COOH; and pharmaceutically acceptable salts thereof:

with the provise that except for the compound wherein R, is \$-CH<sub>a</sub>, R<sub>a</sub> and R<sub>a</sub> taken together form a double bond between positions 9 and 11, R4 and R6 are hydrogen, R42 and R44 taken together form a double bond between positions 4 and 5, Rs is α-F, Ro is β-CHa, Ran is α-OH, Ra and Ras are =0 and Ros is -OP(0)-(OH)o, Ra is =0 only when Roa with Roa forms the above described cyclic phosphate.

Emb-44. The pharmaceutical composition of Emb-43, wherein the angiostatic steroid is selected from the group consisting of: 21-Nor-58-pregnan-9a.17a.20-triol-3-acetate: 21-Nor-5a-pregnan-9a.17a.20-triol-3-phosphate; 21-Nor-58-pregn-17(20)en-3a, 16-diol: 21-Nor-58-pregnan-3a, 176,20-triol: 20-Acetamide-21-nor-58-pregnan-3a, 17α-dipi-3-acetate: 38 Acetamido-58-pregnan-118.17α.21-tripi-20-pne-21-acetate: 21-Nor-5α-pregnan-3α.178. 20-tnol; 21α-Methyl-5β-pregnan-3α,11β, 17α, 21-tetrol-20-one-21-methyl ether; 20-Azido-21-nor-5β-pregnan-3α, 17α-diol; 20(Carbethoxymethyl)thio-21-nor-5β-pregnan-3α,17α-diol; 20-(4-Fluorophenyl)thio-21-nor-5β-pregnan-3α,17α-diol; 16α-(2-Hydroxyethyl)-17β-methyl-5β-androstan-3α,17α-diol; 20-Cyano-21-nor-5β-pregnan-3α,17α-diol; diol: 17a-Methyl-58-endrostan-3a,178-diol: 21-Nor-58-pregn-17(20)en-3a-OL: 21-Nor-58-pregn-17(20)en-3a-ol-S-acetate; 21-Nor-5β-pregn-17(20)-en-3α-ol-16-acetic acid S-acetate; 3β-Azido-5β-pregnan-11β, 17α, 21-triol-20-one-21-acetate; and 5β-Pregnan-11β,17α,21-triol-20-one; 4,9(11)-Pregnadien-17α,21-diol-3,20-dione; 4,9(11) -Pregnedien-17α,21-diol-3,20-dione-21-acetate; 4-Androsten-3-one-17β-carboxylic acid; 17α-Ethynyl-5(10)-estren-17β-ol-3-one; 17α-Ethynyl-1,3,5(10)-estratrien-3,17β-diol; 11-Epicortisol; 17α-Hydroxyprogesterone; Tetrahydrocortexolone; and Tetrahydrocortisol.

Emb-45. The pharmaceutical composition of Emb-44, wherein the anglostatic steroid is selected from the group consisting of: tetrahydrocortisoi; 21-Nor-5β-pregnan-3α,17α,20-triol; 4,9(11)-Pregnadien-17α,21-diol-3,20-dione; 4,9(11)-Pregnadien-17a.21-diol-3,20-dione-21-acetate.

Emb-45. The pharmaceutical composition of Emb- 45 wherein the angiostatic steroid is selected from the group consisting of, tetrahydrocortisol and 4.9(11)-Pregnadien-17g-21-dial-3.20-diane-21-acetate.

Emb-47. A method of treating ophthalmic inflammation without significantly increasing the patient's intraocular pressure, which comprises administering the composition of Emb-43.

Emb-48. A method of treating ophthalmic inflammation without significantly increasing the pateint's intraocular pressure, which comprises: administering the composition of Emb-44.

Emb-49. The method of Emb-49, wherein the angiostatic steroid is selected from the group consisting of: tetrahydrocoriisol; 21-Nor-6}-pregnan-3a,17a,20-triol; 4,9(11)-Pregnadien-17a,21-diol-3,20-dione; 4,9(11)-Pregnadien-17a,21-diol-3,20-dione-21-actate.

# Claims

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#### Use of a compound of the following formula:

Structure [A]

Structure [B]

# wherein

R2

R<sub>1</sub> is H<sub>1</sub>β-CH<sub>2</sub> or β-C<sub>2</sub>H<sub>5</sub>;

R<sub>2</sub> is F, C<sub>9</sub>-C<sub>11</sub> double bond, C<sub>9</sub>-C<sub>11</sub> epoxy, H or -Ct;

is H,  $OR_{28}$ ,  $OC(=O)R_{27}$ , halogen,  $C_{9}C_{11}$  double bond,  $C_{9}C_{11}$  epoxy, =O, OH, -O-silkyf( $C_{1}-C_{12}$ ), -OC ( $=O)Rilkyf(C_{1}-C_{12})$ , -OC ( $=O)Rilkyf(C_{1}-C_{12})$ , -OC ( $=OC)Rilkyf(C_{1}-C_{12})$ , -OC (=OC ), -OC (OC ), -OC

is ARYL as herein defined, or alkyl(C1-C12);

R<sub>4</sub> is H, CH<sub>3</sub>, Cl or F; R<sub>6</sub> is H, OH, F, Cl, Br, CH<sub>3</sub>, phenyl, vinyl or allyl;

R<sub>6</sub> is H or CH<sub>3</sub>;

R<sub>9</sub> is CH<sub>2</sub>CH<sub>2</sub>OR<sub>28</sub>, CH<sub>2</sub>CH<sub>2</sub>OC(=O)R<sub>27</sub>, H, OH, CH<sub>3</sub>, F, =CH<sub>2</sub>, or CH<sub>2</sub>C(=O)OR<sub>28</sub>, OR<sub>28</sub>, O(C=O)R<sub>27</sub>, or

 $\begin{array}{ll} (C=O)CH_2(C=O)OR_{26}; \\ R_{10} & \text{is -C=CH, -CH=CH}_2, \text{ halogen, CN, N}_3, OR_{26}, OC(=O)R_{27}, \text{ H, OH, CH}_3 \text{ or R}_{10} \text{ forms a second bond} \end{array}$ 

between positions C-16 and C-17;

R<sub>12</sub> is H or forms a double bond with R<sub>1</sub> or R<sub>14</sub>; R<sub>13</sub> is halogen, OR<sub>26</sub>, OC(=O)R<sub>27</sub>, NH<sub>2</sub>, NHR<sub>26</sub>, NHC(=O)R<sub>27</sub>, N(R<sub>26</sub>)<sub>2</sub>, NC(=O)R<sub>27</sub>, N<sub>3</sub>, H, -OH, =O, -O-P

 $(=O)(OH)_2$ , or  $=O-C(=O)-(CH)_2(COOH)$  where t is an integer from 2 to 6;  $=H_{14}$  is H or forms a double bond with  $=H_{12}$ ;

R<sub>24</sub> = C, C<sub>1</sub>-C<sub>2</sub> double bond, O;

(R<sub>28</sub>)<sub>2</sub>, that is, an optionally alkyl substituted methylene group; wherein

R<sub>28</sub> = C<sub>1</sub>-C<sub>8</sub> (alkyl, branched alkyl, cycloalkyl, haloalkyl, aralkyl, aryl);

 $R_{27} = R_{26} + OR_{26}$ ;

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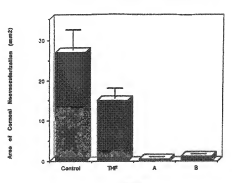
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R<sub>28</sub> = H, C<sub>1</sub>-C<sub>8</sub> (aikyl, branched aikyl, cycloalkyl),

for the preparation of a pharmaceutical composition for preventing and/or treating neovascularization and/or preventing and/or treating ocular hypertension.

- Use according to claim 1, wherein the pharmaceutical composition is for preventing and/or treating ocular neovascularization.
  - 3. Use according to claim 2, wherein the pharmaceutical composition is for preventing and/or treating retinal diseases including including disbetic retinopathy, and some including including disbetic retinopathy, and some including including disbetic retinopathy and some including retinolation are used to subretinal neovascularization; rubeosis titlis; inflammatory diseases; chronic uveitis; neoplasms including retinolatisation and view heteror/bromin indicocyclitis; neovascular glacuscularization; recursing inflammatory, transplantation and developmental hypoplasia of the lifs, neovascularization resulting following a comhised viteratory and lensectory; vascular diseases including retinal ischemia, choroidal vascular insufficiency, choroidal thrombosis and carotid artery ischeriias; pteriglum; neovascularization due to penetration of the oodic neova and neovascularization due to penetration of the oodic neova and neovascularization due to penetration of the oodic neovas and neovascularization due to penetration of the oodic neovas and neovascularization due to penetration of the oodic neovas and neovascularization due to penetration of the oodic neovasculari
  - Use according to claim 1, wherein the pharmaceutical composition is for preventing and/or treating primary open angle glaucoma.

# FIGURE 1 EFFECT OF ANGIOSTATIC STEROIDS ON CORNEAL NEOVASCULARIZATION



Trestment